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Patent application No. Demande de brevet nº

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# PRIORITY DOCUMENT

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Title: A virus causing respiratory tract illness in susceptible mammals.



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The invention relates to the field of virology.

In the past decades several etiological agents of mammalian disease, in particular of respiratory tract illnesses (RTI), in particular of humans, have been identified7. Classical etiological agents of RTI with mammals are respiratory syncytial viruses belonging to the genus *Pneumovirus* found with humans (hRSV) and ruminants such as cattle or sheep (bRSV and/or oRSV). In human RSV differences in reciprocal cross neutralization assays, reactivity of the G proteins in immunological assays and nucleotide sequences of the G gene are used to define 2 hRSV antigenic subgroups. Within the subgroups the aa sequences show 94 % (subgroup A) or 98% (subgroup B) identity, while only 53% aa sequence identity is found between the subgroups. Additional variability is observed within subgroups based on monoclonal antibodies, RT-PCR assays and RNAse protection assays. Viruses from both subgroups have a worldwide distribution and may occur during a single season. Infection may occur in presence of pre-existing immunity and the antigenic variation is not strictly required to allow re-infection. See for example Sullender, W.M., Respiratory Syncytial Virus Genetic and Antigenic Diversity. Clinical Microbiology Reviews, 2000. 13(1): p. 1-15; Collins, P.L., McIntosh, K. and Chanock, R.M., Respiratory syncytial virus. Fields virology, ed. B.N. Knipe, Howley, P.M. 1996, Philadelphia: lippencott-raven. 1313-1351; Johnson, P.R., et al., The G glycoprotein of human respiratory syncytial viruses of subgroups A and B: extensive sequence divergence between antigenically related proteins. Proc Natl Acad Sci U S A, 1987. 84(16): p. 5625-9; Collins, P.L., The molecular Biology of Human Respiratory Syncytial Virus (RSV) of the Genus Pneumovirus, in The Paramyxoviruses, D.W. Kingsbury, Editor. 1991, Plenum Press: New York, p. 103-153.

Another classical *Pneumovirus* is the pneumonia virus of mice (PMV), in general only found with laboratory mice. However, a proportion of the illnesses observed among mammals can still not be attributed to known pathogens.

The invention provides a novel etiological agent, an isolated essentially mammalian negative-sense single stranded RNA virus (herein also called MPV)

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belonging to the subfamily *Pneumovirinae* of the family *Paramyxoviridae* but not identifiable as a classical pneumovirus, and most closely related to and likely belonging to the genus *Metapneumovirus*..

Mammalian viruses resembling metapneumoviruses, i.e. metapneumoviruses isolatable from mammals that essentially function as natural host for said virus or cause disease in said mammals, have until now not been found. Metapneumoviruses, in general thought to be essentially restricted to poultry as natural host or aetiological agent of disease, are also known as avian pneumoviruses. Recently, an AVP isolate of duck was described (FR 2 801 607), further demonstrating that AVP infections are essentially restricted to birds as natural hosts.

The invention provides an isolated mammalian pneumovirus (herein also called MPV) comprising a gene order and amino acid sequence distinct from that of the genus Pneumovirus and which is closely related and considering its phylogenetic relatedness likely belonging to the genus Metapneumovirus within the subfamily Pneumovirinae of the family Paramyxoviridae. Although until now, metapneumoviruses have only been isolated from birds, it is now shown that related, albeit materially distinct, viruses can be identified in other animal species such as mammals. Herein we show repeated isolation of MPV from humans, whereas no such reports exists for APV. Furthermore, unlike APV. MPV essentially does not or only little replicate in chickens and turkeys where it easily does in cynomolgous macaques. No reports have been found on replication of APV in mammals. In addition, whereas specific anti-sera raised against MPV neutralize MPV, anti-sera raised against APV A, B or C do not neutralize MPV, and this lack of cross reactivity provides another proof for MPV being a different metapneumovirus. Furthermore, where APV and MPV share a similar gene order, the G and SH proteins of MPV are largely different from the ones of APV in that they show no significant sequence homologies on both the amino acid or nucleic acid level. Diagnostic assays to discriminate between APV and MPVisolates or antibodies directed against these different viruses can advantageously be developed based on one or both of these proteins (examples are IF, ELISA, neutralization assay, RT-PCR assay). However, also sequence and/or antigenic information obtained from the more related N, P, M, F and L proteins of MPV and analyses of sequence homologies with the respective proteins of APV, can also be used to discriminate between APV and MPV. For

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example, phylogenetic tree analyses of sequence information obtained from MPV again revealed that MPV and APV are two different viruses. In particular, the phylogenetic trees show that APV and MPV are two different lineages of virus. We have also shown that MPV is circulating in the human population for at least 50 years, therefore interspecies transmission has probably taken place at least 50 years ago and is not an everyday event. Since MPV CPE was virtually indistinguishable from that caused by hRSV or hPIV-1 in tMK or other cell cultures, the MPV may have well gone unnoticed until now. The CPE is, as well as with some of the classical *Paramyxoviridae*, characterized by syncytium formation after which the cells showed rapid internal disruption, followed by detachment of the cells from the monolayer. The cells usually (but not always) displayed CPE after three passages of virus from original material, at day 10 to 14 post inoculation, somewhat later than CPE caused by other viruses such as hRSV or hPIV-1.

15 ° Classically, as devastating agents of disease, paramyxoviruses account for many animal and human deaths worldwide each year. The Paramyxoviridae form a family within the order of Mononegavirales (negative-sense single stranded RNA viruses), consisting of the sub-families Paramyxovirinae and Pneumovirinae. The latter sub-family is at present taxonomically divided in the genera Pneumovirus and Metapneumovirus 1. Human respiratory syncytial virus 20 (hRSV), the type species of the Pneumovirus genus, is the single most important cause of lower respiratory tract infections during infancy and early childhood worldwide<sup>2</sup>. Other members of the *Pneumovirus* genus include the bovine and ovine respiratory syncytial viruses and pneumonia virus of mice (PVM). 25 Avian pneumovirus (APV) also known as turkey rhinotracheitis virus (TRTV), the aetiological agent of avian rhinotracheitis, an upper respiratory tract infection of turkeys3, is the sole member of the recently assigned Metapneumovirus genus, which, as said was until now not associated with infections, or what is more, with disease of mammals. Serological subgroups of 30 APV can be differentiated on the basis of sequences of the G glycoprotein and neutralization tests using monoclonal antibodies that also recognize the G glycoprotein, Within subgroups A, B and D the G protein shows 98.5 to 99.7% aa sequence identity within subgroups while between the subgroups only 31.2-38% aa identity is observed. See for example Collins, M.S., Gough, R.E. and

Alexander, D.J., Antigenic differentiation of avian pneumovirus isolates using

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polyclonal antisera and mouse monoclonal antibodies. Avian Pathology, 1993. 22: p. 469-479.; Cook, J.K.A., Jones, B.V., Ellis, M.M., Antigenic differentiation of strains of turkey rhinotracheitis virus using monoclonal antibodies. Avian Pathology, 1993. 22: p. 257-273; Bayon-Auboyer, M.H., et al., Nucleotide sequences of the F, L and G protein genes of two non-A/non-B avian pneumoviruses (APV) reveal a novel APV subgroup. J Gen Virol, 2000. 81(Pt 11): p. 2723-33; Seal, B.S., Matrix protein gene nucleotide and predicted amino acid sequence demonstrate that the first US avian pneumovirus isolate is distinct from European strains. Virus Res, 1998. 58(1-2): p. 45-52; Bayon-Auboyer, M.H., et al., Comparison of F-, G- and N-based RT-PCR protocols with conventional virological procedures for the detection and typing of turkey rhinotracheitis virus. Arch Virol, 1999. 144(6): p. 1091-109; Juhasz, K. and A.J. Easton, Extensive sequence variation in the attachment (G) protein gene of avian pneumovirus: evidence for two distinct subgroups. J Gen Virol, 1994. 75(Pt 11): p. 2873-80.

A further putative serotype of APV is provided in WO00/20600, which describes the Colorado isolate of APV and compared it to known APV or TRT strains with in in vitro serum neutralization tests. First, the Colorado isolate was tested against monospecific polyclonal antisera to recognized TRT isolates. The Colorado isolate was not neutralized by monospecific antisera to any of the TRT strains. It was, however, neutralized by a hyperimmune antiserum raised against a subgroup A strain. This antiserum neutralized the homologous virus to a titre of 1:400 and the Colorado isolate to a titer of 1:80. Using the above method, the Colorado isolate was then tested against TRT monoclonal antibodies. In each case, the reciprocal neutralization titer was <10. Monospecific antiserum raised to the Colorado isolate was also tested against TRT strains of both subgroups. None of the TRT strains tested were neutralized

The Colorado strain of APV does not protect SPF chicks against challenge with either a subgroup A or a subgroup B strain of TRT virus. These results suggest that the Colorado isolate may be the first example of a second serotype of avian pneumovirus, as also suggested by Bayon-Auboyer et al (J. Gen. Vir. 81:2723-2733 (2000).

by the antiserum to the Colorado isolated.

In a preferred embodiment, the invention provides an isolated MPV taxonomically corresponding to a (hereto unknown mammalian) metapneumovirus comprising a gene order distinct from that of the

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pneumoviruses within the sub-familie Pneumovirinae of the family Paramyxoviridae. The classification of the two genera is based primarily on their gene constellation; metapneumoviruses generally lack non-structural proteins such NS1 or NS2 (see also Randhawa et al., J. Vir. 71:9849-9854 (1997) and the gene order is different from that of pneumoviruses (RSV: '3-NS1-NS2-N-P-M-SH-G-F-M2-L-5', APV: '3-N-P-M-F-M2-SH-G-L-5') 4,5,6. MPV as provided by the invention or a virus isolate taxonomically corresponding therewith is upon EM analysis revealed by paramyxovirus-like particles. Consistent with the classification, MPV or virus isolates phylogenetically corresponding or taxonomically corresponding therewith are sensitive to treatment with chloroform; are cultured optimally on tMK cells or cells functionally equivalent thereto and are essentially trypsine dependent in most cell cultures. Furthermore, the typical CPE and lack of haemagglutinating activity with most classically used red blood cells suggested that a virus as provided herein is, albeit only distantly, related to classical pneumoviruses such as RSV. Although most paramyxoviruses have haemagglutinating acitivity, most of the pneumoviruses do not 13. An MPV according to the invention also contains a second overlapping ORF (M2-2) in the nucleic acid fragment encoding the M2 protein, as in general most other pneumoviruses such as for example also demonstrated in Ahmadian et al., J. Gen. Vir. 80:2011-2016 (1999)

To find further viral isolates as provided by the invention it suffices to test a sample, optionally obtained from a diseased animal or human, for the presence of a virus of the sub-family *Pneumovirinae*, and test a thus obtained virus for the presence of genes encoding (functional) NS1 or NS2 or essentially demonstrate a gene order that is different from that of pneumoviruses such as RSV as already discussed above. Furthermore, a virus isolate phylogenetically corresponding and thus taxonomically corresponding with MPV may be found by cross-hybridisation experiments using nucleic acid from a here provided MPV isolate, or in classical cross-serology experiments using monoclonal antibodies specifically directed against and/or antigens and/or immunogens specifically derived from an MPV isolate.

Newly isolated viruses are phylogenetically corresponding to and thus taxonomically corresponding to MPV when comprising a gene order and/or amino acid sequence sufficiently similar to our prototypic MPV isolate(s), or are structurally corresponding therewith, and show close relatedness to the genus

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Metapneumovirus within the subfamily Pneumovirinae. The highest amino sequence homology, and defining the structural correspondence on the individual protein level, between MPV and any of the known other viruses of the same family to date (APV subtype C) is for matrix 87%, for nucleoprotein 88%, for phosphoprotein 68%, for fusionprotein 81% and for parts of the polymerase protein 56-64%%, as can be deduced when comapring the sequences given in figure 6 with sequences of other viruses, in particular of AVP-C. Individual proteins or whole virus isolates with, respectively, higher homology to these mentioned maximum values are considered phylogenetically corresponding and thus taxonomically corresponding to MPV, and comprise a nucleic acid sequence structurally corresponding with a sequence as shown in figure 6a, 6b, 6c.

Herewith the invention provides a virus phylogenetically corresponding to the deposited virus.

It should be noted that, similar to other viruses, a certain degree of variation is found between different isolated essentially mammalian negative-sense single stranded RNA virus isolates as provided herein. In phylogenetic trees, we have identified at least 2 genetic clusters of virus isolates based on comparitive sequence analyses of parts of the L, M, N and F genes. Based on nucleotide and amino-acid differences in the viral nucleic acid or amino acid sequences (the viral sequences), and in analogy to other pneumoviruses such as RSV, these MPV genotypes represent subtypes of MPV. Within each of the genetic clusters of MPV isolates, the percentage identity at the nucleotide level was found to be 94-100 for L, 91-100 for M, 90-100 for N and 93-100 for F and at the amino acid level the percentage identity was found to be 91-100 for L, 98-100 for M, 96-100 for N and 98-100 for F. The minimum percentage identity at the nucleotide level for the entire group of isolated essentially mammalian negative-sense single stranded RNA virus as provided herein (MPV isolates) identified so far was 81 for L and M, 83 for N and 82 for F. At the amino acid level, this percentage was 91 for L and N, 94 for M, and 95 for F. The viral sequence of a MPV isolate or an isolated MPV F gene as provided herein for example shows less than 81% nucleotide sequence identity or less than 82% amino acid sequence identity with the respective nucleotide or amino acid sequence of an APV-C fusion (F) gene as for example provided by Seal et al., Vir. Res. 66: 139147 (2000). Also, the viral sequence of a MPV isolate or an an isolated MPV L gene as provided herein for example shows less than 61% nucleotide sequence identity or less than 63%

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amino acid sequence identity with the respective nucleotide or amino acid sequence of an APV-A polymerase gene as for example provided by Randhawa et al.,, J. Gen. Vir. 77:3047-3051 (1996).

Sequence divergence of MPV strains around the world may be somewhat higher, in analogy with other viruses. Consequently, two potential genetic clusters are identified by analyses of partial nucleotide sequences in the N, M, F and L ORFs of 9 virus isolates. 90-100% nucleotide identity was observed within a cluster, and 81-88% identity was observed between the clusters. Sequence information obtained on more virus isolates confirmed the existence of two genotypes. Virus isolate ned/00/01 as prototype of cluster A, and virus isolate ned/99/01 as prototype of cluster B have been used in cross neutralization assays to test whether the genotypes are related to different serotypes or subgroups.From these data we conclude that essentially mammalian virus isolates displaying percentage amino acid homology higher than 56 for L, 87 for M, 52 for N, 67 for P or 80 for F to isolate I-2614 may definitively be classified as an isolated essentially mammalian negative-sense single stranded RNA virus as provided herein. In particular those virus isolates in general that have a minimum percentage identity at the nucleotide level with a prototype MPV isolate as provided herein of 81 for L and M, 83 for N and/or 82 for F are members of the group of MPV isolates as provided herein . At the amino acid level, these percentage are 91 for L and N, 94 for M, and/or 95 for F. When the percentage amino acid sequence homology for a given virus isolate is higher than 90 for L and N, 93 for M, or 94 for F, the virus isolate is similar to the group of virus isolates displayed in figure 5. When the percentage amino acid sequence homology for a given virus isolate is higher than 94 for L, 95 for N or 97 for M and F the virus isolate can be identified to belong to one of the genotype clusters represented in figure 5. It should be noted that these percentages of homology, by which genetic clusters are defined, are similar to the degree of homology found among genetic clusters in the corresponding genes of RSV.

In short, the invention provides an isolated essentially mammalian negative-sense single stranded RNA virus (MPV) belonging to the sub-familie *Pneumovirinae* of the family *Paramyxoviridae* and identifiable as phylogenetically corresponding to the genus *Metapneumovirus* by determining a nucleic acid sequence of a suitable fragment of the genome of said virus and testing it in phylogenetic tree analyses wherein maximum likelihood trees are

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generated using 100 bootstraps and 3 jumbles and finding it to be more closely phylogenetically corresponding to a virus isolate deposited as I-2614 with CNCM, Paris than it is corresponding to a virus isolate of avian pneumovirus (APV) also known as turkey rhinotracheitis virus (TRTV), the aetiological agent of avian rhinotracheitis.

Suitable nucleic acid genome fragments each useful for such phylogenetic tree analyses are for example any of the RAP-PCR fragments 1 to 10 as disclosed herein in the detailed description, leading to the various phylogenetic tree analyses as disclosed herein in figures 4 or 5. Phylogenetic tree analyses of the nucleoprotein (N), phosphoprotein (P), matrixprotein (M) and fusion protein (F) genes of MPV revealed the highest degree of sequence homology with APV serotype C, the avian pneumovirus found primarily in birds in the United States

In a preferred embodiment, the invention provides an isolated essentially mammalian negative-sense single stranded RNA virus (MPV) belonging to the sub-familie *Pneumovirinae* of the family *Paramyxoviridae* and identifiable as phylogenetically corresponding to the genus *Metapneumovirus* by determining a nucleic acid sequence of a suitable fragment of the genome of said virus and testing it in phylogenetic tree analyses wherein maximum likelihood trees are generated using 100 bootstraps and 3 jumbles and finding it to be more closely phylogenetically corresponding to a virus isolate deposited as I-2614 with CNCM, Paris than it is corresponding to a virus isolate of avian pneumovirus (APV) also known as turkey rhinotracheitis virus (TRTV), the aetiological agent of avian rhinotracheitis, wherein said suitable fragment comprises an open reading frame encoding a viral protein of said virus.

A suitable open reading frame (ORF) comprises the ORF encoding the N protein. When an overall amino acid identity of at least 91%, preferably of at least 95% of the analysed N-protein with the N-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. As shown, the first gene in the genomic map of MPV codes for a 394 amino acid (aa) protein and shows extensive homology with the N protein of other pneumoviruses. The length of the N ORF is identical to the length of the N ORF of APV-C (Table 5) and is smaller than those of other paramyxoviruses (Barr et al., 1991). Analysis of the amino acid sequence revealed the highest homology with APV-C (88%), and only 7-11% with other paramyxoviruses (Table

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Barr et al (1991) identified 3 regions of similarity between viruses belonging to the order Mononegavirales: A, B and C (Figure 8). Although similarities are highest within a virus family, these regions are highly conserved between virus families. In all three regions MPV revealed 97% as sequence identity with APV-C, 89% with APV-B, 92% with APV-A, and 66-73% with RSV and PVM. The region between as residues 160 and 340 appears to be highly conserved among metapneumoviruses and to a somewhat lesser extent the Pneumovirinae (Miyahara et al., 1992; Li et al., 1996; Barr et al., 1991). This is in agreement with MPV being a metapneumovirus, this particular region showing 100% similarity with APV C.

Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the P protein. When an overall amino acid identity of at least 70%, preferably of at least 85% of the analysed P-protein with the P-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. The second ORF in the genome map codes for a 294 aa protein which shares 68% aa sequence homology with the P protein of APV-C, and only 22-26% with the P protein of RSV (Table 6). The P. gene of MPV contains one substantial ORF and in that respect is similar to P from many other paramyxoviruses (Reviewed in Lamb and Kolakofsky, 1996; Sedlmeier et al., 1998). In contrast to APV A and B and PVM and similar to RSV and APV-C the MPV P ORF lacks cysteine residues. Ling (1995) suggested that a region of high similarity between all pneumoviruses (aa 185-241) plays a role in either the RNA synthesis process or in maintaining the structural integrity of the nucleocapsid complex. This region of high similarity is also found in MPV (Figure 9) especially when conservative substitutions are taken in account, showing 100% similarity with APV-C, 93 % with APV-A and B, and approximately 81% with RSV. The C-terminus of the MPV P protein is rich in glutamate residues as has been described for APVs (Ling et al., 1995).

Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the M protein. When an overall amino acid identity of at least 94%, preferably of at least 97% of the analysed M-protein with the M-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. The third ORF of the MPV genome encodes a 254 aa protein, which resembles the M ORFs of other pneumoviruses. The M ORF of MPV has exactly the same size as the M ORFs of

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other metapneumoviruses (Table 5) and shows high as sequence homology with the matrix proteins of APV (78-87%), lower homology with those of RSV and PVM (37-38%) and 10% or less homology with those of other paramyxoviruses (Table 6). Easton (1997) compared the sequences of matrix proteins of all pneumoviruses and found a conserved heptadpeptide at residue 14 to 19 that is also conserved in MPV (Figure 10). For RSV, PVM and APV small secondary ORFs within or overlapping with the major ORF of M have been identified (52 aa and 51 aa in bRSV, 75 aa in RSV, 46 aa in PVM and 51 aa in APV) (Yu et al., 1992; Easton et al., 1997; Samal et al., 1991; Satake et al., 1984). We noticed two small ORFs in the MORF of MPV. One small ORF of 54 aa residues was found within the major M ORF (fragment 1, Figure 7), starting at nucleotide 2281 and one small ORF of 33 aa residues was found overlapping with the major ORF of M starting at nucleotide 2893 (fragment 2, Figure 7). Similar to the secondary ORFs of RSV and APV there is no significant homology between these secondary ORFs and secondary ORFs of the other pneumoviruses, and apparent start or stop signals are lacking. In addition, evidence for the synthesis of proteins corresponding to these secondary ORFs of APV and RSV has not been reported. Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the F protein. When an overall amino acid identity of at least 95%, preferably of at least 97% of the analysed F-protein with the Fprotein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. The F ORF of MPV is located adjacent to the MORF, which is characteristic for members of the Metapneumovirus genus. The F gene of MPV encodes a 539 aa protein, which is two aa residues longer than F of APV-C (Table 5). Analysis of the aa sequence revealed 81% homology with APV-C, 67% with APV-A and B, 33-39% with pneumovirus F proteins and only 10-18% with other paramyxoviruses (Table 6). One of the conserved features among F proteins of paramyxoviruses, and also seen in MPV is the distribution of cysteine residues (Morrison, 1988; Yu et al., 1991). The metapneumoviruses share 12 cysteine residues in F1 (7 are conserved among all paramyxoviruses), and two in F2 (1 is conserved among all paramyxoviruses). Of the 3 potential N-linked glycosylation sites present in the F ORF of MPV, none are shared with RSV and two (position 74 and 389) are shared with APV. The third, unique, potential N-linked glycosylation site for MPV is located at position 206 (Figure 11). Despite the low sequence homology

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with other paramyxoviruses, the F protein of MPV revealed typical fusion protein characteristics consistent with those described for the F proteins of other Paramyxoviridae family members (Morrison, 1988). F proteins of Paramyxoviridae members are synthesized as inactive precursors (F0) that are cleaved by host cell proteases which generate amino terminal F2 subunits and large carboxy terminal F1 subunits. The proposed cleavage site (Collins et al., 1996) is conserved among all members of the Paramyxoviridae family. The cleavage site of MPV contains the residues RQSR. Both arginine (R) residues are shared with APV and RSV, but the glutamine (Q) and serine (S) residues are shared with other paramyxoviruses such as human parainfluenza virus type 1, Sendai virus and morbilliviruses (data not shown). The hydrophobic region at the amino terminus of F1 is thought to function as the membrane fusion domain and shows high sequence similarity among paramyxoviruses and morbilliviruses and to a lesser extend the pneumoviruses (Morrison, 1988). These 26 residues (position 137-163, Figure 11) are conserved between MPV and APV-C, which is in agreement with this region being highly conserved among the metapneumoviruses (Naylor et al., 1998; Seal et al., 2000).

As is seen for the F2 subunits of APV and other paramyxoviruses, MPV revealed a deletion of 22 aa residues compared with RSV (position 107-128, Figure 11). Furthermore, for RSV and APV, the signal peptide and anchor domain were found to be conserved within subtypes and displayed high variability between subtypes (Plows et al., 1995; Naylor et al., 1998). The signal peptide of MPV (aa 10-35, Figure 11) at the amino terminus of F2 exhibits some sequence similarity with APV-C (18 out of 26 aa residues are similar) and less conservation with other APVs or RSV. Much more variability is seen in the membrane anchor domain at the carboxy terminus of F1, although some homology is still seen with APV-C.

Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the M2 protein. When an overall amino acid identity of at least 85%, preferably of at least 90% of the analysed M2-protein with the M2-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. M2 gene is unique to the *Pneumovirinae* and two overlapping ORFs have been observed in all pneumoviruses. The first major ORF represents the M2-1 protein which

enhances the processivity of the viral polymerase (Collins et al., 1995; Collins. 1996) and its readthrough of intergenic regions (Hardy et al., 1998; Fearns et al., 1999). The M2-1 gene for MPV, located adjacent to the F gene, encodes a 187 aa protein (Table 5), and reveals the highest (84%) homology with M2-1 of APV-C 5 (Table 6). Comparison of all pneumovirus M2-1 proteins revealed the highest conservation in the amino-terminal half of the protein (Collins et al., 1990; Zamora et al., 1992; Ahmadian et al., 1999), which is in agreement with the observation that MPV displays 100% similarity with APV-C in the first 80 aa residues of the protein (Figure 12A). The MPV M2-1 protein contains 3 cysteine 10 residues located within the first 30 aa residues that are conserved among all pneumoviruses. Such a concentration of cysteines is frequently found in zincbinding proteins (Ahmadian et al., 1991; Cuesta et al., 2000). The secondary ORFs (M2-2) that overlap with the M2-1 ORFs of pneumoviruses are conserved in location but not in sequence and are thought to be involved in the control of the switch between virus RNA replication and transcription 15 (Collins et al., 1985; Elango et al., 1985; Baybutt et al., 1987; Collins et al., 1990; Ling et al., 1992; Zamora et al., 1992; Alansari et al., 1994; Ahmadian et al., 1999; Bermingham et al., 1999). For MPV, the M2-2 ORF starts at nucleotide 512 in the M2-1 ORF (Figure 7), which is exactly the same start position as for 20 APV-C. The length of the M2-2 ORFs are the same for APV-C and MPV, 71 aa residues (Table 5). Sequence comparison of the M2-2 ORF (Figure 12B) revealed 64% as sequence homology between MPV and APV-C and only 44-48% as sequence homology between MPV and APV-A and B (Table 6). Another suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the L protein. When an overall amino acid identity of at least 91%, preferably of at least 95% of the analysed L-protein with the Lprotein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. In analogy to other negative strand viruses, the last ORF of the MPV genome is the RNA-dependent RNA polymerase component of the replication and transcription complexes. The L 30 gene of MPV encodes a 2005 aa protein, which is 1 residue longer than the APV-A protein (Table 5). The L protein of MPV shares 64% homology with APV-A, 42-44% with RSV, and approximately 13% with other paramyxoviruses (Table 6). Poch et al. (1989; 1990) identified six conserved domains within the L proteins of non-segmented negative strand RNA viruses, from which domain III contained 35

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the four core polymerase motifs that are thought to be essential for polymerase function. These motifs (A, B, C and D) are well conserved in the MPV L protein: in motifs A, B and C: MPV shares 100% similarity with all pneumoviruses and in motif D MPV shares 100% similarity with APV and 92% with RSV's. For the entire domain III (aa 627-903 in the L ORF), MPV shares 77% identity with APV, 61-62% with RSV and 23-27% with other paramyxoviruses (Figure 15). In addition to the polymerase motifs the pneumovirus L proteins contain a sequence which conforms to a consensus ATP binding motif K(X)<sub>21</sub>GEGAGN(X)<sub>20</sub>K (Stec, 1991). The MPV L ORF contains a similar motif as APV, in which the spacing of the intermediate residues is off by one: K(x)<sub>22</sub>GEGAGN(X)<sub>19</sub> K.

A much preferred suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the SH protein. When an overall amino acid identity of at least 30%, preferably of at least 50%, more preferably of at least 75% of the analysed SH-protein with the SH-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. The gene located adjacent to M2 of MPV encodes a 183 aa protein (Figure 7). Analysis of the nucleotide sequence and its deduced amino acid sequence revealed no discernible homology with other RNA virus genes or gene products. The SH ORF of MPV is the longest SH ORF known to date (Table 5). The composition of the aa residues of the SH ORF is relatively similar to that of APV, RSV and PVM, with a high percentage of threonine and serine residues which may represent O-linked glycosylation sites (22%, 18%, 19%, 20.0%, 21% and 28% serine/threonine content for MPV, APV, RSV A, RSV B, bRSV and PVM respectively). The SH ORF of MPV contains 10 cysteine residues, whereas APV SH contains 16 cysteine residues. All pneumoviruses have similar numbers of potential N-glycosylation sites (MPV 2, APV 1, RSV 2, bRSV 3, PVM 4).

The hydrophobicity profiles for the MPV SH protein and SH of APV and RSV revealed similar structural characteristics (Figure 13B). The SH ORFs of APV and MPV have a hydrophylic N-terminus (aa 1-30), a central hydrophobic domain (aa 36-53) which can serve as a potential membrane spanning domain, a second hydrophobic domain around residue 160 and a hydrophilic C-terminus. In contrast, RSV SH appears to lack the C-terminal half of the APV and MPV

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ORFs. In all pneumovirus SH proteins the hydrophobic domain is flanked by basic amino acids, which are also found in the SH ORF for MPV (aa 35 and 54). Another much preferred suitable open reading frame (ORF) useful in phylogenetic analyses comprises the ORF encoding the G protein. When an overall amino acid identity of at least 30%, preferably of at least 50%, more preferably of at least 75% of the analysed G-protein with the G-protein of isolate I-2614 is found, the analysed virus isolate comprises a preferred MPV isolate according to the invention. The G ORF of MPV is located adjacent to the SH gene and encodes a 236 amino acid protein. A secondary small ORF is found immediately following this ORF, potentially coding for 68 aa residues (pos. 6973-7179, fragment 3, Figure 7), but lacking a start codon. A third major ORF, in a different reading frame, of 194 aa residues (fragment 4, Figure 7) is overlapping with both of these ORFs, starting with an ATG codon at nucleotide 6416. This major ORF is followed by a fourth ORF in the same reading frame, possibly coding for 65 aa residues (pos. 7002-7198, fragment 5, Figure 7) but again lacking a start codon. Unlike the first ORF, the other ORFs do not have apparent gene start or gene end sequences (see below). Although the 236 aa residue G ORF probably represents at least a part of the MPV attachment protein it can not be excluded that the additional coding sequences are expressed as separate proteins or as part of the attachment protein through some RNA editing event. It should be noted that for APV and RSV no secondary ORFs after the primary G ORF have been identified but that both APV and RSV have secondary ORFs within the major ORF of G. However, evidence for expression of these ORFs is lacking and there is no homology between the predicted aa sequences for different viruses (Ling et al., 1992). The secondary ORFs in MPV G do not reveal characteristics of other G proteins and whether the additional ORFs are expressed requires further investigation. BLAST analyses with all four ORFs revealed no discernible homology at the nucleotide or aa sequence level with other known virus genes or gene products. This is in agreement with the low sequence homologies found for other G proteins such as hRSV A and B (53%) (Johnson et al., 1987) and APV A and B (38%) (Juhasz et al., 1994). Whereas most of the MPV ORFs resemble those of APV both in length and sequence, the G ORF of MPV is considerably smaller than the G ORF of APV (Table 5). The aa sequence revealed a serine and threonine content of 34%, which is even higher than the 32% for RSV and 24% for APV. The G ORF also contains 8.5% proline

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residues, which is higher than the 8% for RSV and 7% for APV. The unusual abundance of proline residues in the G proteins of APV, RSV and MPV has also been observed in glycoproteins of mucinous origin where it is a major determinant of the proteins three dimensional structure (Collins et al., 1983; Wertz et al., 1985; Jentoft, 1990).

The number of potential N-linked glycosylation sites in G of MPV is similar to other pneumoviruses: MPV has 5, whereas hRSV has 7, bRSV has 5, and APV has 3 to 5.

The predicted hydrophobicity profile of MPV G revealed characteristics similar to the other pneumoviruses. The amino-terminus contains a hydrophylic region followed by a short hydrophobic area (aa 33-53) and a mainly hydrophilic carboxy terminus (Figure 14B). This overall organisation is consistent with that of an anchored type II transmembrane protein and corresponds well with these regions in the G protein of APV and RSV. The G ORF of MPV contains only 1 cysteine residue in contrast to RSV and APV (5 and 20 respectively).

According to classical serological analyses as for example known from Francki, R.I.B., Fauquet, C.M., Knudson, D.L., and Brown, F., Classification and nomenclature of viruses. Fifth report of the international Committee on Taxonomy of Viruses. Arch Virol, 1991. Supplement 2: p. 140-144. an MPV isolate is also identifiable as belonging to a serotype as provided herein, being defined on the basis of its immunological distinctiveness, as determined by quantitative neutralization with animal antisera (obtained from for example ferrets or guinnea pigs as provided in the detailed description). Such a serotype has either no cross-reaction with others or shows a homologous-to heterologous titer ratio >16 in both directions. If neutralization shows a certain degree of cross-reaction between two viruses in either or both directions (homologous-toheterologous tier ration of eight or 16), distinctiveness of serotype is assumed if substantial biophysical/biochemical differences of DNA's exist. If neutralization shows a distinct degree of cross-reaction between two viruses in either or both directions (homologous-to-heterologous tier ration of smaller than eight), identity of serotype of the isolates under study is assumed. As said, useful prototype isolates, such as isolate I-2614, herein also known as MPV isolate 00-1, are provided herein.

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A further classification of a virus as an isolated essentially mammalian negative-sense single stranded RNA virus as provided herein can be made on the basis of homology to the G and/or SH proteins. Where in general the overall amino acid sequence identity between APV (isolated from birds) and MPV (isolated from humans) N, P, M, F, M2 and L ORFs was 64 to 88 percent, and nucleotide sequence homology was also found between the non-coding regions of the APV and MPV genomes, essentially no discernable amino acid sequence homology was found between two of the ORFs of the human isolate (MPV) and any of the ORFs of other paramyxoviruses. The amino acid content, hydrophobicity profiles and location of these ORFs in the viral genome show that they represent G and SH protein analogues. The sequence homology between APV and MPV, their similar genomic organization (3'-N-P-M-F-M2-SH-G-L-5') as well as phylogenetic analyses provide further evidence for the proposed classification of MPV as the first mammalian metapneumovirus. New MPV isolates are for thus example identified as such by virus isolation and characterisation on tMK or other cells, by RT-PCR and/or sequence analysis followed by phylogenetic tree analyses, and by serologic techniques such as virus neutralisation assays, indirect immunofluorescence assays, direct immunofluorescence assays, FACs analyses or other immunological techniques. Preferably these techniques are directed at the SH and/or G protein analogues.

For example the invention provides herein a method to identify further isolates of MPV as provided herein, the method comprising inoculating a essentially MPV-uninfected or specific-pathogen-free guinea pig or ferret (in the detailed description the animal is inoculated intranasally but other ways of inoculation such as intramuscular or intradermal inoculation, and using an other experimental animal, is also feasible) with the prototype isolate I-2614 or related isolates. Sera are collected from the animal at day zero, two weeks and three weeks post inoculation. The animal specifically seroconverted as measured in virus neutralisation (VN) assay and indirect IFA against the respective isolate I-2614 and the sera from the seroconverted animal are used in the immunological detection of said further isolates.

As an example, the invention provides the characterisation of a new member in the family of *Paramyxoviridae*, a human metapneumovirus or metapneumovirus-like virus (since its final taxonomy awaits discussion by a viral taxonomy committee the MPV is herein for example described as

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taxonomically corresponding to APV) (MPV) which may cause severe RTI in humans. The clinical signs of the disease caused by MPV are essentially similar to those caused by hRSV, such as cough, myalgia, vomiting, fever, broncheolitis or pneumonia, possible conjunctivitis, or combinations thereof. As is seen with hRSV infected children, especially very young children may require hospitalisation. As an example an MPV which was deposited January 19, 2001 as I-2614 with CNCM, Institute Pasteur, Paris or a virus isolate phylogenetically corresponding therewith is herewith provided. Therewith, the invention provides a virus comprising a nucleic acid or functional fragment phylogenetically corresponding to a nucleic acid sequence shown in figure 6a, 6b, 6c, or structurally corresponding therewith. In particular the invention provides a virus characterised in that after testing it in phylogenetic tree analyses wherein maximum likelihood trees are generated using 100 bootstraps and 3 jumbles it is found to be more closely phylogenetically corresponding to a virus isolate deposited as I-2614 with CNCM, Paris than it is related to a virus isolate of avian pneumovirus (APV) also known as turkey rhinotracheitis virus (TRTV), the aetiological agent of avian rhinotracheitis.

We propose the new human virus to be named human metapneumovirus or metapneumovirus-like virus (MPV) based on several observations. EM analysis revealed paramyxovirus-like particles. Consistent with the classification, MPV appeared to be sensitive to treatment with chloroform. MPV is cultured optimal on tMK cells and is trypsine dependent. The clinical symptoms caused by MPV as well as the typical CPE and lack of haemagglutinating activity suggested that this virus is closely related to hRSV. Although most paramyxoviruses have haemaglutinating activity, most of the pneumoviruses do not <sup>13</sup>.

As an example, the invention provides a not previously identified paramyxovirus from nasopharyngeal aspirate samples taken from 28 children suffering from severe RTI. The clinical symptoms of these children were largely similar to those caused by hRSV. Twenty-seven of the patients were children below the age of five years and half of these were between 1 and 12 months old. The other patient was 18 years old. All individuals suffered from upper RTI, with symptoms ranging from cough, myalgia, vomiting and fever to broncheolitis and severe pneumonia. The majority of these patients were hospitalised for one

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The virus isolates from these patients had the paramyxovirus morphology in negative contrast electron microscopy but did not react with specific antisera against known human and animal paramyxoviruses. They were all closely related to one another as determined by indirect immunofluorescence assays (IFA) with sera raised against two of the isolates. Sequence analyses of nine of these isolates revealed that the virus is somewhat related to APV. Based on virological data, sequence homology as well as the genomic organisation we propose that the virus is a member of *Metapneumovirus* genus. Serological surveys showed that this virus is a relatively common pathogen since the seroprevalence in the Netherlands approaches 100% of humans by the age of five years. Moreover, the seroprevelance was found to be equally high in sera collected from humans in 1958, indicating this virus has been circulating in the human population for more than 40 years. The identification of this proposed new member of the Metapneumovirus genus now also provides for the development of means and methods for diagnostic assays or test kits and vaccines or serum or antibody compositions for viral respiratory tract infections, and for methods to test or screen for antiviral agents useful in the treatment of MPV infections.

To this extend, the invention provides among others an isolated or recombinant nucleic acid or virus-specific functional fragment thereof obtainable from a virus according to the invention. In particular, the invention provides primers and/or probes suitable for identifying an MPV nucleic acid. Furthermore, the invention provides a vector comprising a nucleic acid according to the invention. To begin with, vectors such as plasmid vectors containing (parts of) the genome of MPV, virus vectors containing (parts of) the genome of MPV. (For example, but not limited to other paramyxoviruses, vaccinia virus, retroviruses, baculovirus), or MPV containing (parts of) the genome of other viruse or other pathogens are provided. Furthermore, a number of techniques have been described for the generation of recombinant negative strand viruses, based on two critical parameters. First, the production of such virus relies on the replication of a partial or full-length copy of the negative sense viral RNA (vRNA) genome or a complementary copy thereof (cRNA). This vRNA or cRNA can be isolated from infectious virus, produced upon in-vitro transcription, or produced in cells upon transfection of nucleic acids. Second, the production of recombinant negative strand virus relies on a functional

polymerase complex. Typically, the polymerase complex of pneumoviruses consists of N, P, L and evt. M2 proteins, but is not necessarily limited thereto. Polymerase complexes or components thereof can be isolated from virus particles, isolated from cells expressing one or more of the components, or 5 produced upon transfection of specific expression vectors. Infectious copies of MPV can be obtained when the above mentioned vRNA. cRNA, or vectors expressing these RNAs are replicated by the above mentioned polymerase complex 16,17,18,19,20,21,22. For the generation of minireplicons or, a reverse genetics system for generating a full-lenght copy comprising most or all 10 of the genome of MPV it suffices to use 3'end and/or 5'end nucleic acid sequences obtainable from for example APV (Randhawa et al., 1997) or MPV itself. Also, the invention provides a host cell comprising a nucleic acid or a vector according to the invention. Plasmid or viral vectors containing the polymerase components of MPV (presumably N, P, L and M2, but not necessarily limited 15 thereto) are generated in prokaryotic cells for the expression of the components in relevant cell types (bacteria, insect cells, eukaryotic cells). Plasmid or viral vectors containing full-length or partial copies of the MPV genome will be generated in prokaryotic cells for the expression of viral nucleic acids in-vitro or in-vivo. The latter vectors may contain other viral sequences for the generation 20 of chimeric viruses or chimeric virus proteins, may lack parts of the viral genome for the generation of replication defective virus, and may contain mutations, deletions or insertions for the generation of attenuated viruses. Infectious copies of MPV (being wild type, attenuated, replication-defective or chimeric) can be produced upon co-expression of the polymerase components 25 according to the state-of-the-art technologies described above. In addition, eukaryotic cells, transiently or stably expressing one or more fulllength or partial MPV proteins can be used. Such cells can be made by transfection (proteins or nucleic acid vectors), infection (viral vectors) or transduction (viral vectors) and may be useful for complementation of mentioned 30 wild type, attenuated, replication-defective or chimeric viruses.

A chimeric virus may be of particular use for the generation of recombinant vaccines protecting against two or more viruses <sup>23,24,26</sup>. For example, it can be envisaged that a MPV virus vector expressing one or more proteins of RSV or a RSV vector expressing one or more proteins of MPV will protect individuals vaccinated with such vector against both virus infections.

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Attenuated and replication-defective viruses may be of use for vaccination purposes with live vaccines as has been suggested for other viruses <sup>25,26</sup>.

In a preferred embodiment, the invention provides a proteinaceous molecule or metapneumovirus-specific viral protein or functional fragment thereof encoded by a nucleic acid according to the invention. Useful proteinaceous molecules are for example derived from any of the genes or genomic fragments derivable from a virus according to the invention. Such molecules, or antigenic fragments thereof, as provided herein, are for example useful in diagnostic methods or kits and in pharmaceutical compositions such as sub-unit vaccines. Particularly useful are the F and/or G protein or antigenic fragments thereof for inclusion as antigen or subunit immunogen, but inactivated whole virus can also be used. Particulary useful are also those proteinaceous subsyances that are encoded by recombinant nucleic acid fragments that are identified for phylogenmetic tree analyses, of course preferred are those that are within the preferred bounds and metes of ORFs useful in phylogenetic tree analyses, in particular for eliciting MPV speciofic antibodies, be it in vivo (e.g. for protective puposes or for providing diagnostic antibodies) or in vitro (e.g. by phage display technology or another technique useful for generating synthetic antibodies).

Also provided herein are antibodies, be it natural polyclonal or monoclonal, or synthetic (e.g. (phage) library-derived binding molecules) antibodies that specifically react with an antigen comprising a proteinaceous molecule or MPV-specific functional fragment thereof according to the invention. Such antibodies are useful in a method for identifying a viral isolate as an MPV comprising reacting said viral isolate or a component thereof with an antibody as provided herein. This can for example be achieved by using purified or non-purified MPV or parts thereof (proteins, peptides) using ELISA, RIA, FACS or similar formats of antigen detection assays (Current Protocols in Immunology). Alternatively, infected cells or cell cultures may be used to identify viral antigens using classical immunofluorescence or immunohistochemical techniques.

Other methods for identifying a viral isolate as a MPV comprise reacting said viral isolate or a component thereof with a virus specific nucleic acid according to the invention, in particular where said mammalian virus comprises a human virus.

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In this way the invention provides a viral isolate identifiable with a method according to the invention as a mammalian virus taxonomically corresponding to a negative-sense single stranded RNA virus identifiable as likely belonging to the genus *Metapneumovirus* within the sub-familie *Pneumoviriae* of the family *Paramyxoviriae*.

The method is useful in a method for virologically diagnosing an MPV infection of a mammal, said method for example comprising determining in a sample of said mammal the presence of a viral isolate or component thereof by reacting said sample with a nucleic acid or an antibody according to the invention. Examples are further given in the detailed description, such as the use of PCR (or other amplification or hybridisation techniques well known in the art) or the use of immunofluorescence detection (or other immunological techniques known in the art)

The invention also provides a method for serologically diagnosing a MPV infection of a mammal comprising determining in a sample of said mammal the presence of an antibody specially directed against a MPV or component thereof by reacting said sample with a proteinaceous molecule or fragment thereof or an antigen according to the invention.

Methods and means provided herein are particularly useful in a diagnostic kit for diagnosing a MPV infection, be it by virological or serological diagnosis. Such kits or assays may for example comprise a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an antigen and/or an antibody according to the invention.

Use of a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an antigen and/or an antibody according to the invention is also provided for the production of a pharmaceutical composition, for example for the treatment or prevention of MPV infections and/or for the treatment or prevention of respiratory tract illnesses, in particular in humans. Attenuation of the virus can be achieved by established methods developed for this purpose, including but not limited to the use of related viruses of other species, serial passages through laboratory animals or/and tissue/cell cultures, site directed mutagenesis of molecular clones and exchange of genes or gene fragments between related viruses.

A pharmaceutical composition comprising a virus, a nucleic acid, a proteinaceous molecule or fragment thereof, an antigen and/or an antibody

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according to the invention can for example be used in a method for the treatment or prevention of a MPV infection and/or a respiratory illness comprising providing an individual with a pharmaceutical composition according to the invention. This is most useful when said individual comprises a human, especially when said human is below 5 years of age, since such infants and young children are most likely to be infected by a human MPV as provided herein. Generally, in the acute phase patients will suffer from upper respiratory symptoms predisposing for other respiratory and other diseases. Also lower respiratory illnesses may occur, predisposing for more and other serious conditions.

The invention also provides method to obtain an antiviral agent useful in the treatment of respiratory tract illness comprising establishing a cell culture or experimental animal comprising a virus according to any one of claims 1 to 7, treating said culture or animal with an candidate antiviral agent, and determining the effect of said agent on said virus or its infection of said culture or animal. An example of such an antiviral agent comprises a MPV-neutralising antibody, or functional component thereof, as provided herein, but antiviral agents of other nature are obtained as well. The invention also provides use of an antiviral agent according to the invention for the preparation of a pharmaceutical composition, in particular for the preparation of a pharmaceutical composition for the treatment of respiratory tract illness. especially when caused by an MPV infection, and provides a pharmaceutical composition comprising an antiviral agent according to the invention, useful in a method for the treatment or prevention of an MPV infection or respiratory illness, said method comprising providing an individual with such a pharmaceutical composition.

The invention is further explained in the detailed description without limiting it thereto.

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### Figure legends

Figure 1A comprises table 1: Percentage homology found between the amino acid sequence of isolate 00-1 and other members of the Pneumovirinae. Percentages (x100) are given for the amino acid sequences of N, P, M, F and two RAP-PCR fragments in L (8 and 9/10). Accession numbers used for the analyses are described in the materials and methods section.

Fig 1B comprises table 2: Seroprevalence of MPV in humans categorised by age group using immunofluorescence and virus neutralisation assays..

Fig. 2: Schematic representation of the genome of APV with the location and size of the fragments obtained with RAP-PCR and RT-PCR on virus isolate 00-1. Fragments 1 to 10 were obtained using RAP-PCR. Fragment A was obtained with a primer in RAP-PCR fragment 1 and 2 and a primer designed based on alignment of leader and trailer sequences of APV and RSV<sup>6</sup>. Fragment B was obtained using primers designed in RAP-PCR fragment 1 and 2 and RAP-PCR fragment 3. Fragment C was obtained with primers designed in RAP-PCR fragment 3 and RAP-PCR fragment 4,5,6 and 7.

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For all phylogenetic trees, (figures 3-5) DNA sequences were aligned using the ClustalW software package and maximum likelihood trees were generated using the DNA-ML software package of the Phylip 3.5 program using 100 bootstraps and 3 jumbles <sup>15</sup>. Previously published sequences that were used for the generation of phylogenetic trees are available from Genbank under accessions numbers: For all ORFs: hRSV: NC001781; bRSV: NC001989; For the FORF: PVM, D11128; APV-A, D00850; APV-B, Y14292; APV-C, AF187152; For the NORF: PVM, D10331; APV-A, U39295; APV-B, U39296; APV-C, AF176590; For the MORF: PMV, U66893; APV-A, X58639; APV-B, U37586; APV-C, AF262571; For the PORF: PVM, 09649; APV-A, U22110, APV-C, AF176591. Phylogenetic

For the P ORF: PVM, 09649; APV-A, U22110, APV-C, AF176591. Phylogenetic analyses for the nine different virus isolates of MPV were performed with APV strain C as outgroup.

Abbreviations used in figures: hRSV: human RSV; bRSV: bovine RSV; PVM: pneumonia virus of mice; APV-A,B, and C: avian pneumovirus typa A, B and C.

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Fig. 3 Comparison of the N, P, M and F ORF's of members of the subfamily *Pneumovirinae* and virus isolate 00-1. The alignment shows the amino acid sequence of the complete N, P, M and F proteins and partial L proteins of virus isolate 00-1. Amino acids that differ between isolate 00-1 and the other viruses are shown, identical amino acids are represented by periods, gaps are represented as dashes. Numbers correspond to amino acid positions in the proteins. Accession numbers used for the analyses are described in the materials and methods section. APV-A, B or C: Avian Pneumovirus type A, B or C, b-or hRSV: bovine or human respiratory syncytial virus, PVM: pneumonia virus of mice. L8: fragment 8 obtained with RAP-PCR located in L, L9/10: consensus of fragment 9 and 10 obtained with RAP-PCR, located in L. For the P allignment, no APV-B sequence was available from the Genebank, For the L allignment only bRSV, hRSV and APV-A sequences were available.

Fig. 4: Phylogenetic analyses of the N, P, M, and F ORF's of members of the genus Pneumovirinae and virus isolate 00-1. Phylogenetic analysis was performed on viral sequences from the following genes: F (panel A), N (panel B), M (panel C), and P (panel D). The phylogenetic trees are based on maximum likelyhood analyses using 100 bootstraps and 3 jumbles. The scale representing the number of nucleotide changes is shown for each tree.

Fig. 5: Phylogenetic relationship for parts of the F (panel A), N (panel B), M (panel C) and L (panel D) ORFs of nine of the primary MPV isolates with APV-C, it's closest relative genetically. The phylogenetic trees are based on maximum likelyhood analyses. The scale representing the number of nucleotide changes is shown for each tree. Accession numbers for APV-C: panel A: D00850; panel B: U39295; panel C: X58639; and panel D: U65312.

Fig. 6A: Nucleotide and amino acid sequence information from the 3'end of the genome of MPV isolate 00-1.ORF's are given. N: ORF for nucleoprotein; P: ORF for phosphoprotein; M: ORF for matrix protein; F: ORF for fusion protein; GE: gene end; GS: gene start.

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Fig. 6B and C: Nucleotide and amino acid sequence information from obtained fragments in the polymerase gene (L) of MPV isolates 00-1. Positioning of the fragments in L is based on protein homologies with APV-C (accession number U65312). The translated fragment 8 (Fig. 6B.) is located at amino acid number 8 to 243, and the consensus of fragments 9 and 10 (Fig. 6C) is located at amino acid number 1358 to 1464 of the APV-C L ORF.

# Figure 7

Genomic map of MPV isolate 00-1. The nucleotide positions of the start and stop codons are indicated under each ORF. Unknown secondary ORFs longer than 50 aa residues are indicated by numbers and placed above the primary ORFs. The double lines which cross the L ORF indicate the shortened representation of the L gene.

# Figure 8:

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Alignment of the predicted amino acid sequence of the nucleoprotein of MPV with those of other pneumoviruses. The conserved regions identified by Barr (1991) are represented by boxes and labelled A, B, and C. The conserved region among pneumoviruses (Li, 1996) is shown gray shaded. Gaps are represented by dashes, periods indicate the positions of identical amino acid residues compared to MPV.

#### Figure 9:

Amino acid sequence comparison of the phosphoprotein of MPV with those of other pneumoviruses. The region of high similarity (Ling, 1995) is boxed, and the glutamate rich region is grey shaded. Gaps are represented by dashes and periods indicate the position of identical amino acid residues compared to MPV.

# Figure 10:

Comparison of the deduced amino acid sequence of the matrix protein of MPV with those of other pneumoviruses. The conserved heptad sequence (Easton, 1997) is grey

shaded. Gaps are represented by dashes and periods indicate the position of identical amino acid residues relative to MPV.

#### Figure 11:

Allignment of the predicted amino acid sequence of the fusion protein of MPV with those of other pneumoviruses. The conserved cysteine residues are boxed, N-linked glycosylation sites are underlined, the cleavage site of F0 is double underlined, the fusion peptide, signal peptide and membrane anchor domain are shown grey shaded. Gaps are represented by dashes and periods indicate the position of identical amino acids relative to MPV.

#### Figure 12

Comparison of amino acid sequence of the M2 ORFs of MPV with those of other pneumoviruses. The alignment of M2-1 ORFs is shown in panel A, with the conserved amino terminus (Collins, 1990; Zamora, 1999) shown grey shaded. The three conserved cysteine residues are printed bold face and indicated by #. The alignment of M2-2 ORFs is shown in panel B. Gaps are represented by dashes and periods indicate the position of identical amino acids relative to MPV.

# 20 Figure 13

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Amino acid sequence analyses of the SH ORF of MPV. (A) Amino acid sequence of the SH ORF of MPV, with the serine and threonine residues grey shaded, cysteine residues in bold face and the hydrophobic region double underlined. Potential N-linked glycosylation sites are single underlined. Numbers indicate the positions of the basic amino acids flanking the hydrophobic domain.(B) Alignment of the hydrophobicity plots of the SH proteins of MPV, APV-A and hRSV-B. The procedure of Kyte and Doolittle (1982) was used with a window of 17 amino acids. Arrows indicate a strong hydrophobic domain. Positions within the ORF are given on the X-axis.

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#### Figure 14

Amino acid sequence analyses of the G ORF of MPV. (A) Amino acid sequence of the G ORF of MPV, with serine, threonine and proline residues grey shaded, the cysteine residue is in bold face and the hydrophobic region double underlined. The potential N-linked glycosylation sites are single underlined. (B) Alingment of the

hydrophobicity plots of the G proteins of MPV, APV-A and hRSV-B. The procedure of Kyte and Doolittle (1982) was used with a window of 17 amino acids. Arrows indicate the hydrophobic region, and positions within the ORF are given at the X-axis.

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# Figure 15

Comparison of the amino acid sequences of a conserved domain of the polymerase gene of MPV and other paramyxoviruses. Domain III is shown with the four conserved polymerase motifs (A, B, C, D) in domain III (Poch 1998, 1999) boxed. Gaps are represented by dashes and periods indicate the position of identical amino acid residues relative to MPV. hPIV3: human parainfluenza virus type 3; SV: sendai virus; hPIV-2: human parainfluenza virus type 2; NDV: New castle disease virus; MV: measles virus; PDV: phocine distemper virus; nipah: Nipah virus.

# 15 Figure 16:

Phylogenetic analyses of the M2-1 and L ORFs of MPV and selected paramyxoviruses. The M2-1 ORF was aligned with the M2-1 ORFs of other members of the genus *Pneumovirinae* (A) and the L ORF was aligned with L ORFs members of the genus *pneumovirinae* and selected other paramyxoviruses as described in the legends of figure 15 (B). Phylogenetic trees were generated by maximum likelihood analyses using 100 bootstraps and 3 jumbles. The scale representing the number of nucleotide changes is shown for each tree. Numbers in the trees represent bootstrap values based on the consensus trees.

#### 25 Figure 17:

Noncoding sequences of MPV isolate 00-1. (A) The gene junctions, with sequences shown from 3' to 5'end in the viral RNA. Each segment represents a gene junction, containing the end of a gene, followed by the number of additional nucleotides in the intergenic region and the start signal for the next gene. Initiation and stop codons are underlined. Panel B shows the nucleotide sequence of the 3'leader and 5'trailer regions of the MPV genome. The leader and trailer sequences of MPV are aligned with each other, and with the leader and trailer sequences of APV. Underlined regions represent the primer sequences used in RT-PCR assays which are based on the 3' and 5' end sequences of APV and RSV (Randhawa et al., 1997; Mink et al.,

1991). The gray shaded regions indicate part of the start signal for the N ORF. Le: leader, Tr. trailer.

Detailed description

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Virus isolation and characterisation

From 1980 till 2000 we found 28 unidentified virus isolates from patients with severe respiratory disease. These 28 unidentified virus isolates grew slowly in tMK cells, poorly in VERO cells and A549 cells and could not or only little be propagated in MDCK or chicken embryonated fibroblast cells. Most of these virus isolates induced CPE after three passages on tMK cells, between day ten and fourteen. The CPE was virtually indistinguishable from that caused by hRSV or hPIV in tMK or other cell cultures, characterised by syncytium formation after which the cells showed rapid internal disruption, followed by detachment of the cells from the monolayer. The cells usually (sometimes later) displayed CPE after three passages of virus from original material, at day 10 to 14 post inoculation, somewhat later than CPE caused by other viruses such as hRSV or hPIV.

We used the supernatants of infected tMK cells for EM analysis which revealed the presence of paramyxovirus-like virus particles ranging from 150 to 600 nanometer, with short envelope projections ranging from 13 to 17 nanaometer. Consistent with the biochemical properties of enveloped viruses such as the *Paramyxoviridae*, standard chloroform or ether treatment<sup>8</sup> resulted in >10<sup>4</sup> TCID50 reduction of infectivity for tMK cells. Virus-infected tMK cell culture supernatants did not display heamagglutinating activity with turkey, chicken and guinea pig erythrocytes. During culture, the virus replication appeared to be trypsine dependent on the cells tested. These combined virological data allowed that the newly identified virus was taxonomically classified as a member of the *Paramyxoviridae* family.

We isolated RNA from tMK cells infected with 15 of the unidentified virus isolates for reverse transcription and polymerase chain reaction (RT-PCR) analyses using primer-sets specific for *Paramyxovirinae*<sup>9</sup>, hPIV 1-4, sendai virus, simian virus type 5, New-Castle disease virus, hRSV, morbilli, mumps, Nipah, Hendra, Tupaia and Mapuera viruses. RT-PCR assays were carried out at low stringency in order to detect potentially related viruses and RNA isolated from

homologous virus stocks were used as controls. Whereas the available controls reacted positive with the respective virus-specific primers, the newly identified virus isolates did not react with any primer set, indicating the virus was not closely related to the viruses tested.

We used two of the virus-infected tMK cell culture supernatants to inoculate guinea pigs and ferrets intranasaly. Sera were collected from these animals at day zero, two weeks and three weeks post inoculation. The animals displayed no clinical symptoms but all seroconverted as measured in virus neutralisation (VN) assays and indirect IFA against the homologous viruses. The sera did not react in indirect IFA with any of the known paramyxoviruses described above and with PVM. Next, we screened the so far unidentified virus isolates using the guinea pig and ferret pre- and post-infection sera, of which 28 were clearly positive by indirect IFA with the post-infection sera suggesting they were serological closely related or identical.

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#### RAP PCR

To obtain sequence information on the unknown virus isolates, we used a random PCR amplification strategy known as RAP-PCR10. To this end, tMK cells were infected with one of the virus isolates (isolate 00-1) as well as with hPIV-1 which served as a control. After both cultures displayed similar levels of CPE, virus in the culture supernatants was purified on continuous 20-60% sucrose gradients. The gradient fractions were inspected for virus-like particles by EM, and RNA was isolated from the fraction containing approximately 50% sucrose, in which nucleocapsids were observed. Equivalent amounts of RNA isolated from both virus fractions were used for RAP-PCR, after which samples were run side by side on a 3% NuSieve agarose gel. Twenty differentially displayed bands specific for the unidentified virus were subsequently purified from the gel, cloned in plasmid pCR2.1 (Invitrogen) and sequenced with vector-specific primers. When we used these sequences to search for homologies against sequences in the Genbank database using the BLAST software (www.ncbi.nlm.nih.gov/BLAST/) 10 out of 20 fragments displayed resemblance to APV/TRTV sequences. These 10 fragments were located in the genes coding for the nucleoprotein (N; fragment 1 and 2), the matrix protein (M; fragment 3), the fusion protein (F; fragment 4, 5, 6, 7,) and the polymerase protein (L; fragment 8,9,10) (Fig.2). We next designed PCR primers to complete the sequence information for the 3' end

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of the viral genome based on our RAP PCR fragments as well as published leader and trailer sequences for the Pneumovirinae 6. Three fragments were amplified, of which fragment A spanned the extreme 3' end of the N open reading frame (ORF), fragment B spanned the phosphoprotein (P) ORF and fragment C closed the gap between the M and F ORFs (Fig. 2). Sequence analyses of these three fragments revealed the absence of NS1 and NS2 ORFs at the extreme 3' end of the viral genome and positioning of the FORF immediately adjacent to the M ORF. This genomic organisation resembles that of the metapneumovirus APV. which is also consistent with the sequence homology. Overall the translated sequences for the N, P, M and F ORFs showed an average of 30% homology with members of the genus Pneumovirus and 66% with members of the genus Metapneumovirus. The amino acid homologies found for N showed about 40% homology with hRSV and 88% with APV-C, its closest relative genetically, as for example can be deduced by comparing the amino acid sequence of figure 3 with the amino acid sequence of the respective N proteins of other viruses. The amino acid sequence for P showed about 25% homology with hRSV and about 67% with APV-C, M showed about 37% with hRSV and about 87% with APV-C, F showed about 40% homology with hRSV and about 81% with APV-C, and the fragments obtained in L showed an average of 33% with hRSV and 55% with APV-C.

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#### Phylogeny

Although BLAST searches using nucleotide sequences obtained from the unidentified virus isolate revealed homologies primarily with members of the Pneumovirinae, homologies based on protein sequences revealed some resemblance with other paramyxoviruses as well (data not shown). As an indication for the relation between the newly identified virus isolate and members of the *Pneumovirinae*, phylogenetic trees were constructed based on the N, P, M and F ORFs of these viruses. In all four phylogenetic trees, the newly identified virus isolate was most closely related to APV (Fig.4). From the four serotypes of APV that have been described<sup>11</sup>, APV serotype C, the metapneumovirus found primarily in birds in the USA, showed the closest resemblance to the newly identified virus. It should be noted however, that only partial sequence information for APV serotype D is available.

To determine the relationship of our various newly identified virus isolates, we constructed phylogenetic trees based on sequence information

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obtained from eight to nine isolates (8 for F, 9 for N, M and L). To this end, we used RT-PCR with primers designed to amplify short fragments in the N, M, F and L ORFs, that were subsequently sequenced directly. The nine virus isolates that were previously found to be related in serological terms (see above) were also found to be closely related genetically. In fact, all nine isolates were more closely related to one another than to APV. Although the sequence information used for these phylogenetic trees was limited, it appears that the nine isolates can be divided in two groups, with isolate 94-1, 99-1 and 99-2 clustering in one group and the other six isolates (94-2; 93-1; 93-2; 93-3; 93-4; 00-1) in the other (Fig.5).

### Seroprevalence

To study the seroprevalence of this virus in the human population, we tested sera from humans in different age categories by indirect IFA using tMK cells infected with one of the unidentified virus isolates. This analysis revealed that 25% of the children between six and twelve months had antibodies to the virus, and by the age of five nearly 100% of the children were seropositive. In total 56 serum samples tested by indirect IFA were tested by VN assay. For 51 (91%) of the samples the results of the VN assay (titre >8) coincided with the results obtained with indirect IFA (titre>32). Four samples that were found positive in IFA, were negative by VN test (titre <8) whereas one serum reacted negative in IFA (titre<32) and positive in the VN test (titre 16) (table 2). IFA conducted with 72 sera taken from humans in 1958 (ages ranging from 8-99 years)<sup>12,27</sup> revealed a 100% seroprevalence, indicating the virus has been circulating in the human population for more than 40 years. In addition a number of these sera were used in VN assays to confirm the IFA data (table 2).

Genetic analyses of the N, M, P and F genes revealed that MPV has higher sequence homology to the recently proposed genus *Metapneumovirinae* (average of 63 %) as compared to the genus *Pneumovirinae* (average of 30 %) and thus demonstrates a genomic organisation similar to and resembling that of APV/TRTV. In contrast to the genomic organisation of the RSVs ('3-NS1-NS2-N-P-M-SH-G-F-M2-L-5'), metapneumoviruses lack NS1 and NS2 genes and have a different positioning of the genes between M and L ('3-N-P-M-F-M2-SH-G-L-5').

35 The lack of ORFs between the M and F genes in our virus isolates and the lack of

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NS1 and NS2 adjacent to to N, and the high amino acid sequence homology found with APV are reasons to propose the classification of MPV isolated from humans as a first member of the *Metapneumovirus* genus of mammalian, in particular of human origin.

Phylogenetic analyses revealed that the nine MPV isolates from which sequence information was obtained are closely related. Although sequence information was limited, they were in fact more closely related to one another than to any of the avian metapneumoviruses. Of the four serotypes of APV that have been described, serotype C was most closely related to MPV based on the N. P, M and F genes. It should be noted however that for serotype D only partial sequences for the F gene were available from Genbank and for serotype B only M, N and F sequences were available. Our MPV isolates formed two clusters in phylogenetic trees. For both hRSV and APV different genetic and serological subtypes have been described. Whether the two genetic clusters of MPV isolates represent serogical subgroups that are also functionally different remains unknown at presentOur serological surveys showed that MPV is a common human pathogen. The repeated isolation of this virus from clinical samples from children with severe RTI indicates that the clinical and economical impact of MPV may be high. New diagnostic assays based on virus detection and serology will allow a more detailed analysis of the incidence and clinical and economical impact of this viral pathogen.

The slight differences between the IFA and VN results (5 samples) maybe due to the fact that in the IFA only IgG serum antibodies were detected whereas the VN assay detects both classes and sub-classes of antibodies or differences may be due to the differences in sensitivity between both assays. For IFA a cut off value of 16 is used, whereas for VN a cut off value of 8 is used.

On the other hand, differences between IFA versus VN assay may also indicate possible differences between different serotypes of this newly identified virus. Since MPV seems most closely related to APV, we speculate that the human virus may have originated from birds. Analysis of serum samples taken from humans in 1958 revealed that MPV has been widespread in the human population for more then 40 years indicating that a tentative zoonosis event must have taken place long before 1958.

#### Materials and Methods

# Specimen collection

5 Over the past decades our laboratory has collected nasopharyngeal aspirates from children suffering from RTI, which are routinely tested for the presence of viruses. All nasopharyngeal aspirates were tested by direct immmunofluorescence assays (DIF) using fluorescence labelled antibodies against influenza virus types A, and B, hRSV and human parainfluenza virus 10 (hPIV) types 1 to 3. The nasopharyngeal aspirates were also processed for virus isolation using rapid shell vial techniques 14 on various celllines including VERO cells, tertiary cynomolgous monkey kidney (tMK) cells, human endothelial lung (HEL) cells and marbin dock kidney (MDCK) cells. Samples showing cytophatic effects (CPE) after two to three passages, and which were negative in DIF, were 15 tested by indirect immunofluorescence assays (IFA) using virus specific antibodies against influenza virus types A, B and C, hRSV types A and B, measles virus, mumps virus, human parainfluenza virus (hPIV) types 1 to 4, sendai virus, simian virus type 5, and New-Castle disease virus. Although for many cases the aetiological agent could be identified, some specimens were negative for all these viruses tested. 20

#### Direct Immunofluorescence Assay (DIF)

Nasopharyngeal aspirate samples from patients suffering from RTI were used for DIF and virus isolation as described 14,15. Samples were stored at -70 °C. In brief, nasopharyngeal aspirates were diluted with 5 ml Dulbecco MEM (BioWhittaker, Walkersville, MD) and thoroughly mixed on a vortex mixer for one minute. The suspension was thus centrifuged for ten minutes at 840 x g. The sediment was spread on a multispot slide (Nutacon, Leimuiden, The Netherlands), the supernatant was used for virus isolation. After drying, the cells were fixed in aceton for 1 minute at room temperature. After washing the slides were incubated for 15 minutes at 37 °C with commercial available FITC-labelled virus specific anti-sera such as influenza A and B, hRSV and hPIV 1 to 3 (Dako, Glostrup, Denmark). After three washings in PBS and one in tap water, the slides were included in a glycerol/PBS solution (Citifluor, UKC,

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Canterbury, UK) and covered. The slides were analysed using a Axioscop fluorescence microscope (Carl Zeiss B.V, Weesp, the Netherlands.

#### Virus isolation

For virus isolation tMK cells (RIVM, Bilthoven, The Netherlands) were cultured 5 in 24 well plates containing glass slides (Costar, Cambridge, UK), with the medium described below supplemented with 10% fetal bovine serum (BioWhittaker, Vervier, Belgium). Before inoculation the plates were washed with PBS and supplied with Eagle's MEM with Hanks' salt (ICN, Costa mesa, 10 CA) of which half a litre was supplemented with 0.26 gram HaHCO<sub>3</sub>, 0.025 M Hepes (Biowhittaker), 2 mM L-glutamine (Biowhittaker), 100 units penicilline, 100 µg streptomycine (Biowhittaker), 0.5 gram lactalbumine (Sigma-Aldrich, Zwijndrecht, The Netherlands), 1.0 gram D-glucose (Merck, Amsterdam, The Netherlands), 5.0 gram peptone (Oxoid, Haarlem, The Netherlands) and 0.02% 15 trypsine (Life Technologies, Bethesda, MD). The plates were inoculated with supernatant of the nasopharyngeal aspirate samples, 0,2 ml per well in triplicate, followed by centrifuging at 840x g for one hour. After inoculation the plates were incubated at 37 °C for a maximum of 14 days changing the medium once a week and cultures were checked daily for CPE. After 14 days cells were 20 scraped from the second passage and incubated 14 days. This step was repeated for the third passage. The glass slides were used to demonstrate the presence of the virus by indirect IFA as described below.

#### Animal immunisation

Ferret and guinea pig specific antisera for the newly discovered virus were generated by experimental intranasal infection of two specific pathogen free ferrets and two guinea pigs, housed in separate pressurised glove boxes. Two to three weeks later all the animals were bled by cardiac puncture, and their sera were used as reference sera. The sera were tested for all previous described viruses with indirect IFA as described below.

# Antigen detection by indirect IFA

We performed indirect IFA on slides containing infected tMK cells. After washing with PBS the slides were incubated for 30 minutes at 37 °C with virus specific anti-sera. We used monoclonal antibodies in DIF against influenza A, B

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and C, hPIV type 1 to 3 and hRSV as described above. For hPIV type 4, mumps virus, measles virus, sendai virus, simian virus type 5, New-Castle Disease virus polyclonal antibodies (RIVM) and ferret and guinea pig reference sera were used. After three washings with PBS and one wash with tap water, the slides were stained with a secondary antibodies directed against the sera used in the first incubation. Secondary antibodies for the polyclonal anti sera were goat-antiferret (KPL, Guilford, UK, 40 fold diluted), mouse-anti-rabbit (Dako, Glostrup, Denmark, 20 fold diluted), rabbit-anti-chicken (KPL, 20 fold dilution) and mouse-anti-guinea pig (Dako, 20 fold diluted). Slides were processed as described for DIF.

# Detection of antibodies in humans by indirect IFA

For the detection of virus specific antibodies, infected tMK cells were fixed with cold acetone on coverslips, washed with PBS and stained with serum samples at a 1 to 16 dilution. Subsequently, samples were stained with FITC-labelled rabbit anti human antibodies 80 times diluted in PBS (Dako). Slides were processed as described above.

# Virus culture of MPV

Sub-confluent mono-layers of tMK cells in media as described above were inoculated with supernatants of samples that displayed CPE after two or three passages in the 24 well plates. Cultures were checked for CPE daily and the media was changed once a week. Since CPE differed for each isolate, all cultures were tested at day 12 to 14 with indirect IFA using ferret antibodies against the new virus isolate. Positive cultures were freeze-thawed three times, after which the supernatants were clarified by low-speed centrifugation, aliquoted and stored frozen at -70 °C. The 50% tissue culture infectious doses (TCID50) of virus in the culture supernatants were determined as described.

#### 30 Virus neutralisation assay

VN assays were performed with serial two-fold dilutions of human and animal sera starting at an eight-fold dilution. Diluted sera were incubated for one hour with 100 TCID50 of virus before inoculation of tMK cells grown in 96 well plates, after which the plates were centrifuged at 840 x g. The media was changed after three and six days and IFA was conducted with ferret antibodies against MPV 8

days after inoculation. The VN titre was defined as the lowest dilution of the serum sample resulting in negative IFA and inhibition of CPE in cell cultures.

#### Virus characterisation

- Haemagglutination assays and chloroform sensitivity tests were performed as described<sup>8,14</sup>. For EM analyses, virus was concentrated from infected cell culture supernatants in a micro-centrifuge at 4 °C at 17000 x g, after which the pellet was resuspended in PBS and inspected by negative contrast EM. For RAP-PCR, virus was concentrated from infected tMK cell supernatants by ultra-
- centrifugation on a 60% sucrose cussion (2 hours at 150000 x g, 4 °C). The 60% sucrose interphase was subsequently diluted with PBS and layered on top of a 20-60% continuous sucrose gradient which was centrifuged for 16 hours at 275000 x g at 4 °C. Sucrose gradient fractions were inspected for the presence of virus-like particles by EM and poly-acrylamide gel electrophoresis followed by silver staining. The approximately 50% sucrose fractions that appeared to contain nucleocapsids were used for RNA isolation and RAP-PCR.

#### RNA isolation

RNA was isolated from the supernatant of infected cell cultures or sucrose

20 gradient fractions using a High Pure RNA Isolation kit according to instructions
from the manufacturer (Roche Diagnostics, Almere, The Netherlands).

#### RT-PCR

- Virus-specific oligonucleotide sequences for RT-PCR assays on known
  paramyxoviruses are described in addenda 1. A one-step RT-PCR was performed
  in 50 µl reactions containing 50 mM Tris.HCl pH 8.5, 50 mM NaCl, 4 mM MgCl<sub>2</sub>,
  2 mM dithiotreitol, 200 µM each dNTP, 10 units recombinant RNAsin (Promega,
  Leiden, the Netherlands), 10 units AMV RT (Promega, Leiden, The
  Netherlands), 5 units Amplitaq Gold DNA polymerase (PE Biosystems,
- Nieuwerkerk aan de Ijssel, The Netherlands) and 5 μl RNA. Cycling conditions were 45 min. at 42 °C and 7 min. at 95 °C once, 1 min at 95 °C, 2 min. at 42 °C and 3 min. at 72 °C repeated 40 times and 10 min. at 72 °C once.



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#### RAP-PCR

RAP-PCR was performed essentially as described10. The oligonucleotide sequences are described in addenda 2. For the RT reaction, 2 µl RNA was used in a 10  $\mu$ l reaction containing 10 ng/ $\mu$ l oligonucleotide, 10 mM dithiotreitol, 500  $\mu$ m 5 each dNTP, 25 mM Tris-HCl pH 8.3, 75 mM KCl and 3 mM MgCl<sub>2</sub>. The reaction mixture was incubated for 5 min. at 70 °C and 5 min. at 37 °C, after which 200 units Superscript RT enzyme (LifeTechnologies) were added. The incubation at 37 °C was continued for 55 min. and the reaction terminated by a 5 min. incubation at 72 °C. The RT mixture was diluted to give a 50 µl PCR reaction 10 containing 8 ng/µl oligonucleotide, 300 µm each dNTP, 15 mM Tris-HCL pH 8.3. 65 mM KCl, 3.0 mM MgCL2 and 5 units Taq DNA polymerase (PE Biosystems). Cycling conditions were 5 min. at 94 °C, 5 min. at 40 °C and 1 min. at 72 °C once, followed by 1 min. at 94 °C, 2 min. at 56 °C and 1 min. at 72 °C repeated 40 15 times and 5 min. at 72°C once. After RAP-PCR, 15 µl the RT-PCR products were run side by side on a 3% NuSieve agarose gel (FMC BioProducts, Heerhugowaard, The Netherlands). Differentially displayed fragments specific for MPV were purified from the gel with Qiaquick Gel Extraction kit (Qiagen. Leusden, The Netherlands) and cloned in pCR2.1 vector (Invitrogen, Groningen,

### Sequence analysis

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RAP-PCR products cloned in vector pCR2.1 (Invitrogen) were sequenced with M13-specific oligonucleotides. DNA fragments obtained by RT-PCR were purified from agarose gels using Qiaquick Gel Extraction kit (Qiagen, Leusden, The Netherlands), and sequenced directly with the same oligonucleotides used for PCR. Sequence analyses were performed using a Dyenamic ET terminator sequencing kit (Amersham Pharmacia Biotech, Roosendaal, The Netherlands) and an ABI 373 automatic DNA sequencer (PE Biosystem). All techniques were performed according to the instructions of the manufacturer.

The Netherlands) according to instructions from the manufacturer.

# Generating genomic fragments of MPV by RT-PCR

To generate PCR fragments spanning gaps A, B and C between the RAP-PCR fragments (Fig.2) we used RT-PCR assays as described before on RNA isolated

5 from virus isolate 00-1. The following primers were used:
For fragment A: TR1 designed in the leader: (5'AAAGAATTCACGAGAAAAAAACGC-3') and N1 designed at the 3'end of the
RAP-PCR fragments obtained in N (5'-CTGTGGTCTCTAGTCCCACTTC-3')
For fragment B: N2 designed at the 5'end of the RAP-PCR fragments obtained in
N: (5'-CATGCAAGCTTATGGGGC-3') and M1 designed at the 3'end of the RAPPCR fragments obtained in M: (5'-CAGAGTGGTTATTGTCAGGGT-3').
For fragment C: M2 designed at the 5'end of the RAP-PCR fragment obtained in
M: (5'-GTAGAACTAGGAGCATATG-3') and F1 designed at the 3'end of the RAP-

15 Fragments were purified from the gel, cloned and sequenced as described before.

PCR fragments obtained in F: (5'-TCCCCAATGTAGATACTGCTTC-3').

RT-PCR for diagnosing MPV.

For the amplification and sequencing of parts of the N, M, F and L ORFs of nine
of the MPV isolates, we used primers N3 (5'-GCACTCAAGAGATACCCTAG -3')
and N4 (5'-AGACTTTCTGCTTTGCTGCCTG-3'), amplifying a 151 nucleotide
fragments, M3 (5'-CCCTGACAATAACCACTCTG-3') and M4 (5'GCCAACTGATTTGGCTGAGCTC-3') amplifying a 252 nucleotide fragment, F7
(5'-TGCACTATCTCCTCTTGGGGGCTTTG-3') and F8 (5'-

TCAAAGCTGCTTGACACTGGCC-3') amplifying a 221 nucleotide fragment and L6 (5'-CATGCCCACTATAAAAGGTCAG-3') and L7 (5'-CACCCCAGTCTTTCTTGAAA-3') amplifying a 173 nucleotide fragment respectively. RT-PCR, gel purification and direct sequencing were performed as described above. Furthermore, probes used were:

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Probe used in M: 5'-TGC TTG TAC TTC CCA AAG-3'
Probe used in N: 5'-TAT TTG AAC AAA AAG TGT-3'
Probe used in L: 5'-TGGTGTGGGATATTAACAG-3'

## Phylogenetic analyses

For all phylogenetic trees, DNA sequences were alligned using the ClustalW software package and maximum likelihood trees were generated using the DNA-

- ML software package of the Phylip 3.5 program using 100 bootstraps and 3 jumbles<sup>15</sup>. Previously published sequences that were used for the generation of phylogenetic trees are available from Genbank under accessions numbers: For all ORFs: hRSV: NC001781; bRSV: NC001989; For the F ORF: PVM, D11128; APV-A, D00850; APV-B, Y14292; APV-C, AF187152; For the N ORF: PVM,
- D10331; APV-A, U39295; APV-B, U39296; APV-C, AF176590; For the M ORF: PMV,U66893; APV-A, X58639; APV-B, U37586; APV-C, AF262571; For the P ORF: PVM, 09649; APV-A, U22110, APV-C, AF176591. Phylogenetic analyses for the nine different virus isolates of MPV were performed with APV strain C as outgroup.

Abbreviations used in figures: hRSV: human RSV; bRSV: bovine RSV; PVM: pneumonia virus of mice; APV-A, B, and C: avian pneumovirus typ A, B and C.

# Examples of methods to identify MPV

20 Specimen collection

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In order to find virus isolates nasopharyngeal aspirates, throat and nasal swabs, broncheo alveolar lavages preferably from mammals such as humans, carnivors (dogs, cats, mustellits, seals etc.), horses, ruminants (cattle, sheep, goats etc.), pigs, rabbits, birds (poultry, ostridges, etc) should be examined. From birds cloaca swabs and droppings can be examined as well. Sera should be collected for immunological assays, such as ELISA and virus neutralisation assays. Collected virus specimens were diluted with 5 ml Dulbecco MEM medium (BioWhittaker, Walkersville, MD) and thoroughly mixed on a vortex mixer for one minute. The suspension was thus centrifuged for ten minutes at 840 x g. The sediment was spread on a multispot slide (Nutacon, Leimuiden, The Netherlands) for immunofluorescence techniques, and the supernatant was used for virus isolation.

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#### Virus isolation

For virus isolation tMK cells (RIVM, Bilthoven, The Netherlands) were cultured in 24 well plates containing glass slides (Costar, Cambridge, UK), with the medium described below supplemented with 10% fetal bovine serum 5 (BioWhittaker, Vervier, Belgium). Before inoculation the plates were washed with PBS and supplied with Eagle's MEM with Hanks' salt (ICN, Costa mesa, CA) supplemented with 0.52/liter gram NaHCOs, 0.025 M Hepes (Biowhittaker), 2 mM L-glutamine (Biowhittaker), 200 units/liter penicilline, 200 µg/liter streptomycine (Biowhittaker), 1gram/liter lactalbumine (Sigma-Aldrich, 10 Zwijndrecht, The Netherlands), 2.0 gram/liter D-glucose (Merck, Amsterdam, The Netherlands), 10 gram/liter peptone (Oxoid, Haarlem, The Netherlands) and 0.02% trypsine (Life Technologies, Bethesda, MD). The plates were inoculated with supernatant of the nasopharyngeal aspirate samples, 0,2 ml per well in 15 triplicate, followed by centrifuging at 840x g for one hour. After inoculation the plates were incubated at 37 °C for a maximum of 14 days changing the medium once a week and cultures were checked daily for CPE. After 14 days, cells were scraped from the second passage and incubated for another 14 days. This step was repeated for the third passage. The glass slides were used to demonstrate the presence of the virus by indirect IFA as described below. 20 CPE was generally observed after the third passage, at day 8 to 14 depending on the isolate. The CPE was virtually indistinghuisable from that caused by hRSV or hPIV in tMK or other cell cultures. However, hRSV induces CPE starting around day 4. CPE was characterised by syncytia formation, after which the cells showed rapid internal disruption, followed by detachment of cells from the 25 monolayer. For some isolates CPE was difficult to observe, and IFA was used to confirm the presence of the virus in these cultures.

#### Virus culture of MPV

30 Sub-confluent monolayers of tMK cells in media as described above were inoculated with supernatants of samples that displayed CPE after two or three passages in the 24 well plates. Cultures were checked for CPE daily and the media was changed once a week. Since CPE differed for each isolate, all cultures were tested at day 12 to 14 with indirect IFA using ferret antibodies against the new virus isolate. Positive cultures were freeze-thawed three times, after which

the supernatants were clarified by low-speed centrifugation, aliquoted and stored frozen at -70 °C. The 50% tissue culture infectious doses (TCID50) of virus in the culture supernatants were determined following established techniques used in the field<sup>16</sup>.

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#### Virus characterisation

Haemagglutination assays and chloroform sensitivity tests were performed following well established and described techniques used in the field<sup>14</sup>. For EM analyses, virus was concentrated from infected cell culture supernatants in a micro-centrifuge at 4 °C at 17000 x g, after which the pellet was resuspended in PBS and inspected by negative contrast EM.

## Antigen detection by indirect IFA

Weesp, the Netherlands).

Collected specimens were processed as described and sediment of the samples was spread on a multispot slide. After drying, the cells were fixed in aceton for 1 minute at room temperature.

Alternatively, virus was cultured on tMK cells in 24 well slides containing glass slides. These glass slides were washed with PBS and fixed in aceton for 1 minute at room temperature.

After washing with PBS the slides were incubated for 30 minutes at 37 °C with polyclonal antibodies at a dilution of 1:50 to 1:100 in PBS. We used immunised ferrets and guinea pigs to obtain polyclonal antibodies, but these antibodies can be raised in various animals, and the working dilution of the polyclonal antibody can vary for each immunisation. After three washes with PBS and one wash with tap water, the slides were incubated at 37°C for 30 minutes with FITC labeled goat-anti-ferret antibodies (KPL, Guilford, UK, 40 fold diluted). After three washes in PBS and one in tap water, the slides were included in a glycerol/PBS solution (Citifluor, UKC, Canterbury, UK) and covered. The slides were analysed using an Axioscop fluorescence microscope (Carl Zeiss B.V.,

Detection of antibodies in humans, mammals, ruminants or other animals by indirect IFA

For the detection of virus specific antibodies, infected tMK cells with MPV were fixed with acetone on coverslips (as described above), washed with PBS and incubated 30 minutes at 37°C with serum samples at a 1 to 16 dilution. After two washes with PBS and one with tap water, the slides were incubated 30 minutes at 37°C with FITC-labelled secondary antibodies to the species used (Dako). Slides were processed as described above.

Antibodies can be labelled directly with a fluorescent dye, which will result in a direct immuno fluorescence assay. FITC can be replaced with any fluorescent dye.

#### Animal immunisation

Ferret and guinea pig specific antisera for the newly discovered virus were generated by experimental intranasal infection of two specific pathogen free ferrets and two guinea pigs, housed in separate pressurised glove boxes. Two to three weeks later the animals were bled by cardiac puncture, and their sera were used as reference sera. The sera were tested for all previous described viruses with indirect IFA as described below. Other animal species are also suitable for the generation of specific antibody preparations and other antigen preparations may be used.

## Virus neutralisation assay (VN assay)

VN assays were performed with serial two-fold dilutions of human and animal sera starting at an eight-fold dilution. Diluted sera were incubated for one hour with 100 TCID50 of virus before inoculation of tMK cells grown in 96 well plates, after which the plates were centrifuged at 840 x g. The same culture media as described above was used. The media was changed after three and six days, and after 8 days IFA was performed (see above). The VN titre was defined as the lowest dilution of the serum sample resulting in negative IFA and inhibition of CPE in cell cultures.

#### RNA isolation

RNA was isolated from the supernatant of infected cell cultures or sucrose gradient fractions using a High Pure RNA Isolation kit according to instructions from the manufacturer (Roche Diagnostics, Almere, The Netherlands). RNA can also be isolated following other procedures known in the field (Current Protocols in Molecular Biology).

## 10 RT-PCR

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A one-step RT-PCR was performed in 50 μl reactions containing 50 mM Tris.HCl pH 8.5, 50 mM NaCl, 4 mM MgCl<sub>2</sub>, 2 mM dithiotreitol, 200 μM each dNTP, 10 units recombinant RNAsin (Promega, Leiden, the Netherlands), 10 units AMV RT (Promega, Leiden, The Netherlands), 5 units Amplitaq Gold DNA polymerase (PE Biosystems, Nieuwerkerk aan de Ijssel, The Netherlands) and 5 μl RNA. Cycling conditions were 45 min. at 42 °C and 7 min. at 95 °C once, 1 min at 95 °C, 2 min. at 42 °C and 3 min. at 72 °C repeated 40 times and 10 min. at 72 °C once.

Primers used for diagnostic PCR:

In the nucleoprotein: N3 (5'-GCACTCAAGAGATACCCTAG -3') and N4 (5'-AGACTTTCTGCTTGCTGCCTG-3'), amplifying a 151 nucleotide fragment.

In the matrixprotein: M3 (5'-CCCTGACAATAACCACTCTG-3') and M4 (5'-GCCAACTGATTTGGCTGAGCTC-3') amplifying a 252 nucleotide fragment.

In the polymerase protein: L6 (5'-CATGCCCACTATAAAAGGTCAG-3') and L7

(5'-CACCCCAGTCTTTCTTGAAA-3') amplifying a 173 nucleotide fragment.

Other primers can be designed based on MPV sequences, and different buffers and assay conditions may be used for specific purposes.

#### Sequence analysis

Sequence analyses were performed using a Dyenamic ET terminator sequencing kit (Amersham Pharmacia Biotech, Roosendaal, The Netherlands) and an ABI 373 automatic DNA sequencer (PE Biosystem). All techniques were performed according to the instructions of the manufacturer. PCR fragments were sequenced directly with the same oligonucleotides used for PCR, or the fragments were purified from the gel with Qiaquick Gel Extraction kit (Qiagen,

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Leusden, The Netherlands) and cloned in pCR2.1 vector (Invitrogen, Groningen, The Netherlands) according to instructions from the manufacturer and subsequently sequenced with M13-specific oligonucleotides.

Oligonucleotides used for analysing the 3'end of the genome (absence of 5 NS1/NS2).

Primer TR1 (5'-AAAGAATTCACGAGAAAAAAACGC-3') was designed based on published sequences of the trailer and leader for hRSV and APV, published by Randhawa (1997) and primer N1 (5'-CTGTGGTCTCTAGTCCCACTTC-3') was

designed based on obtained sequences in the N protein. The RT-PCR assay and sequencing was performed as described above.

The RT-PCR gave a product of approximately 500 base pairs which is to small to contain information for two ORFS, and translation of these sequences did not reveal an ORF.

Detection of antibodies in humans, mammals, ruminants or other animals by **ELISA** 

In Paramyxoviridae, the N protein is the most abundant protein, and the immune response to this protein occurs early in infection. For these reasons, a recombinant source of the N proteins is preferably used for developing an ELISA assay for detection of antibodies to MPV. Antigens suitable for antibody detection include any MPV protein that combines with any MPV-specific antibody of a patient exposed to or infected with MPV virus. Preferred antigens of the invention include those that predominantly engender the immune 25 response in patients exposed to MPV, which therefore, typically are recognised most readily by antibodies of a patient. Particularly preferred antigens include the N, F and G proteins of MPV.

Antigens used for immunological techniques can be native antigens or can be modified versions thereof. Well known techniques of molecular biology can be used to alter the amino acid sequence of a MPV antigen to produce modified versions of the antigen that may be used in immunologic techniques. Methods for cloning genes, for manipulating the genes to and from expression vectors, and for expressing the protein encoded by the gene in a heterologous host are well-known, and these techniques can be used to provide the expression

vectors, host cells, and the for expressing cloned genes encoding antigens in a host to produce recombinant antigens for use in diagnostic assays. See for instance: Molecular cloning, A laboratory manual and Current Protocols in Molecular Biology.

- A variety of expression systems may be used to produce MPV antigens. For instance, a variety of expression vectors suitable to produce proteins in *E.Coli*, *B.subtilis*, yeast, insect cells and mammalian cells have been described, any of which might be used to produce a MPV antigen suitable to detect anti-MPV antibodies in exposed patients.
- The baculovirus expression system has the advantage of providing necessary processing of proteins, and is therefor preferred. The system utilizes the polyhedrin promoter to direct expression of MPV antigens. (Matsuura et al. 1987, J.Gen.Virol. 68: 1233-1250).
- Antigens produced by recombinant baculo-viruses can be used in a variety of immunological assays to detect anti-MPV antibodies in a patient. It is well established, that recombinant antigens can be used in place of natural virus in practically any immunological assay for detection of virus specific antibodies. The assays include direct and indirect assays, sandwich assays, solid phase assays such as those using plates or beads among others, and liquid phase assays. Assays suitable include those that use primary and secondary antibodies, and those that use antibody binding reagents such as protein A. Moreover, a variety of detection methods can be used in the invention, including colorimetric, fluorescent, phosphorescent, chemiluminescent, luminescent and radioactive methods.

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Example 1 of indirect anti-PMV IgG EIA using recombinant N protein

An indirect IgG EIA using a recombinant N protein (produced with recombinant baculo-virus in insect (Sf9) cells) as antigen can be performed. For antigen preparation, Sf9 cells are infected with the recombinant baculovirus and harvested 3-7 days post infection. The cell suspension is washed twice in PBS, pH 7.2, adjusted to a cell density of 5.0X 10<sup>6</sup> cells/ml, and freeze-thawed three times. Large cellular debris is pelleted by low speed centrifugation (500 x g for 15 min.) and the supernatant is collected and stored at -70°C until use. Uninfected cells are processed similarly for negative control antigen.

100 µl of a freeze-thaw lysate is used to coat microtiter plates, at dilutions ranging from 1:50 to 1:1000. An uninfected cell lysate is run in duplicate wells and serves as a negative control. After incubation overnight, plates are washed twice with PBS/0.05%Tween. Test sera are diluted 1:50 to 1:200 in ELISA buffer (PBS, supplemented to 2% with normal goat sera, and with 0.5% bovine serum albumine and 0.1% milk), followed by incubation wells for 1 hour at 37°C.

Plates are washed two times with PBS/0.05%Tween. Horseradish peroxidase labelled goat anti-human (or against other species) IgG, diluted 1:3000 to 1:5000 in ELISA buffer, added to wells, and incubated for 1 hour at 37°. The plates are then washed two times with PBS/0.05%Tween and once with tap water, incubated for 15 minutes at room temperature with the enzyme substrate TMB, 3,3',5,5' tetramethylbenzidine, such as that obtained from Sigma, and the reaction is stopped with 100 µl of 2 M phosphoric acid. Colorimetric readings are measured at 450 nm using an automated microtiter plate reader.

# Example 2: Capture anti-MPV IgM EIA using a recombinant nucleoprotein

- A capture IgM EIA using the recombinant nucleoprotein or any other recombinant protein as antigen can be performed by modification of assays as previously described by Erdman et al (1990) J.Clin.Microb. 29: 1466-1471.

  Affinity purified anti-human IgM capture antibody (or against other species), such as that obtained from Dako, is added to wells of a microtiter plate in a concentration of 250 ng per well in 0.1 M carbonate buffer pH 9.6. After overnight incubation at room temperature, the plates are washed two times with PBS/0.05% Tween. 100 μl of test serum diluted 1:200 to 1:1000 in ELISA buffer is added to triplicate wells and incubated for 1 hour at 37°C. The plates are then washed two times with in PBS/0.05%Tween.
- The freeze-thawed (infected with recombinant virus) Sf21 cell lysate is diluted 1:100 to 1: 500 in ELISA buffer is added to the wells and incubated for 2 hours at 37°C. Uninfected cell lysate serves as a negative control and is run in duplicate wells.
  - The plates are then washed three times in PBS/0.05% Tween and incubated for 1 hour at 37°C with 100 µl of a polyclonal antibody against MPV in a optimal

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dilution in ELISA buffer. After 2 washes with PBS/0.05% Tween, the plates are incubated with horseradish peroxide labeled secondary antibody (such as rabbit anti ferret), and the plates are incubated 20 minutes at 37°C.

The plates are then washed five times in PBS/0/05% Tween, incubated for 15 minutes at room temperature with the enzyme substrate TMB, 3,3',5,5' tetramethylbenzidine, as, for instance obtained from "Sigma", and the reaction is stopped with 100 µl of 2M phosphoric acid. Colormetric readings are measured at 450 nm using automated microtiter plate reader.

The sensitivities of the capture IgM EIAs using the recombinant nucleoprotein (or other recombinant protein) and whole MPV virus are compared using acuteand convalescent-phase serum pairs form persons with clinical MPV virus infection. The specificity of the recombinant nucleoprotein capture EIA is determined by testing serum specimens from healthy persons and persons with other paramyxovirus infections.

Potential for EIAs for using recombinant MPV fusion and glycoprotein proteins produced by the baculovirus expression.

The glycoproteins G and F are the two transmembraneous envelope glycoproteins of the MPV virion and represent the major neutralisation and protective antigens.

The expression of these glycoproteins in a vector virus system sych as a baculovirus system provides a source of recombinant antigens for use in assays for detection of MPV specific antibodies. Moreover, their use in combination with the nucleoprotein, for instance, further enhances the sensitivity of enzyme immunoassays in the detection of antibodies against MPV.

A variety of other immunological assays (Current Protocols in Immunology) may

be used as alternative methods to those described here.

In order to find virus isolates nasopharyngeal aspirates, throat and nasal swabs,
broncheo alveolar lavages and throat swabs preferable from but not limited to
humans, carnivors (dogs, cats, mustellits, seals etc.), horses, ruminants (cattle,
sheep, goats etc.), pigs, rabbits, birds (poultry, ostridges, etc) can be examined.

From birds classe and intestinal and the state of the state o

35 From birds, cloaca and intestinal swabs and droppings can be examined as well.

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For all samples, serology (antibody and antigen detection etc.), virus isolation and nucleic acid detection techniques can be performed for the detection of virus. Monoclonal antibodies can be generated by immunising mice (or other animals) with purified MPV or parts thereof (proteins, peptides) and subsequently using established hybridoma technology (Current protocols in Immunology).

Alternatively, phage display technology can be used for this purpose (Current protocols in Immunology).

Similarly, polyclonal antibodies can be obtained from infected humans or animals, or from immunised humans or animals (Current protocols in Immunology).

The detection of the presence or absence of NS1 and NS2 proteins can be performed using western-blotting, IFA, immuno precipitation techniques using a variety of antibody preparations. The detection of the presence or absence of NS1 and NS2 genes or homologues thereof in virus isolates can be performed using PCR with primer sets designed on the basis of known NS1 and/or NS2 genes as well as with a variety of nucleic acid hybridisation techniques.

To determine whether NS1 and NS2 genes are present at the 3' end of the viral genome, a PCR can be performed with primers specific for this 3' end of the genome. In our case, we used a primer specific for the 3' untranslated region of the viral genome and a primer in the N ORF. Other primers may be designed for the same purpose. The absence of the NS1/NS2 genes is revealed by the length and/or nucleotide sequence of the PCR product. Primers specific for NS1 and/or NS2 genes may be used in combination with primers specific for other parts of the 3' end of the viral genome (such as the untranslated region or N, M or F ORFs) to allow a positive identification of the presence of NS1 or NS2 genes. In addition to PCR, a variety of techniques such as molecular cloning, nucleic acid hybridisation may be used for the same purpose.

# Example 3: Different serotypes/subgroups of MPV

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Two potential genetic clusters are identified by analyses of partial nucleotide sequences in the N, M, F and L ORFs of 9 virus isolates. 90 -100% nucleotide identity was observed within a cluster, and 81-88% identity was observed between the clusters. Sequence information obtained on more virus isolates

confirmed the existence of two genotypes. Virus isolate ned/00/01 as prototype of cluster A, and virus isolate ned/99/01 as prototype of cluster B have been used in cross neutralization assays to test whether the genotypes are related to different serotypes or subgroups.

## Results

Using RT-PCR assays with primers located in the polymerase gene, we identified 30 additional virus isolates from nasopharyngeal aspirate samples. Sequence information of parts of the matrix and polymerase genes of these new isolates together with those of the previous 9 isolates were used to construct phylogenetic trees (Figure 16). Analyses of these trees confirmed the presence of two genetic clusters, with virus isolate ned/00/00-1 as the prototype virus in group A and virus isolate ned/99/01 as the prototype virus in group B. The nucleotide sequence identity within a group was more than 92%, while between the clusters the identity was 81-85%.

Virus isolates ned/00/01 and ned/99/01 have been used to inoculate ferrets to raise virus-specific antisera. These antisera were used in virus neutralization assays with both viruses.

Table 3: Virus neutralization titers

	isolate 00-1	isolate 99-1
preserum	□2	□2
ferret A		
(00-1)		
ferret A	64	□2
22 dpi		
(00-1)		
preserum	□2	□ 2
ferret B		
(99-1)		
ferret B	4	64
22 dpi		
(99-1)		

For isolate 00-1 the titer differs 32 (64/2) fold

5 For isolate 99-1 the titer differs 16 (64/4) fold

In addition, 6 guinea pigs have been inoculated with either one of the viruses (ned/00/01 and ned/99/01). RT-PCR assays on nasopharyngeal aspirate samples showed virus replication from day 2 till day 10 post infection. At day 70 post infection the guinea pigs have been challenged with either the homologous or the heterologous virus, and for in all four cases virus replication has been noticed.

Table 4

	primary	virus	secondary	virus	
	infection	replication	infection	replication	
guinea pig 1-3	00-1	2 out of 3	99-1	1 out of 2	
guinea pig 4-6	00-1	3 out of 3	00-1	1 out of 3	
guinea pig 7-9	99-1	3 out of 3	00-1	2 out of 2	
guinea pig 10-12	99-1	3 out of 3	99-1	1 out of 3	

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note: for the secondary infection guinea pig 2 and 9 were not there any more.

Virus neutralization assays with anti sera after the first challenge showed essentially the same results as in the VN assays performed with the ferrets (> 16-fold difference in VN titer).

The results presented in this example confirm the existence of two genotypes, which correspond to two serotypes of MPV, and show the possibility of repeated infection with heterologous and homologous virus

## Example 4: Further sequence determination

This example describes the further analysis of the sequences of MPV open reading frames (ORFs) and intergenic sequences as well as partial sequences of the genomic termini.

Sequence analyses of the nucleoprotein (N), phosphoprotein (P), matrixprotein (M) and fusion protein (F) genes of MPV revealed the highest degree of sequence homology with APV serotype C, the avian pneumovirus found primarily in birds in the United States. These analyses also revealed the absence of non-structural proteins NS1 and NS2 at the 3'end of the viral genome and positioning of the fusion protein immediately adjacent to the matrix protein. Here we present the sequences of the 22K (M2) protein, the small hydrophobic (SH) protein, the attachment (G) protein and the polymerase (L) protein genes, the intergenic regions and the trailer sequence. In combination with the sequences described previously the sequences presented here complete the genomic sequence of MPV with the exception of the extreme 12-15 nucleotides of the genomic termini and establish the genomic organisation of MPV. Side by side comparisons of the sequences of the MPV genome with those of APV subtype A, B and C, RSV subtype A and B, PVM and other paramyxoviruses provides strong evidence for the classification of MPV in the *Metapneumovirus* genus.

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#### Results

# Sequence strategy

MPV isolate 00-1 (van den Hoogen et al., 2001) was propagated in tertiary monkey kidney (tMK) cells and RNA isolated from the supernatant 3 weeks after inoculation was used as template for RT-PCR analyses. Primers were designed on the basis of the partial sequence information available for MPV 00-1 (van den Hoogen et al., 2001) as well as the leader and trailer sequences of APV and RSV (Randhawa et al., 1997; Mink et al., 1991). Initially, fragments between the previously obtained products, ranging in size from 500 bp to 4 Kb in length, were generated by RT-PCR amplification and sequenced directly. The genomic sequence was subsequently confirmed by generating a series of overlapping RT-PCR fragments ranging in size from 500 to 800 bp that represented the entire MPV genome. For all PCR fragments, both strands were sequenced directly to minimize amplification and sequencing errors. The nucleotide and amino acid sequences were used to search for homologies with sequences in the Genbank database using the BLAST software (www.ncbi.nlm.nih.gov/BLAST). protein names were assigned to open reading frames (ORFs) based on homology with known viral genes as well as their location in the genome. Based on this information, a genomic map for MPV was constructed (Figure 7). The MPV genome is 13378 nucleotides in length and its organization is similar to the genomic organization of APV. Below, we present a comparison between the ORFs and non-coding sequences of MPV and those of other paramyxoviruses and discuss the important similarities and differences.

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#### The nucleoprotein (N) gene

As shown, the first gene in the genomic map of MPV codes for a 394 amino acid (aa) protein and shows extensive homology with the N protein of other pneumoviruses. The length of the N ORF is identical to the length of the N ORF of APV-C (Table 5) and is smaller than those of other paramyxoviruses (Barr et al., 1991). Analysis of the amino acid sequence revealed the highest homology with APV-C (88%), and only 7-11% with other paramyxoviruses (Table 6). Barr et al (1991) identified 3 regions of similarity between viruses belonging to the order Mononegavirales: A, B and C (Figure 8). Although similarities are highest within a virus family, these regions are highly conserved between virus

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families. In all three regions MPV revealed 97% aa sequence identity with APV-C, 89% with APV-B, 92% with APV-A, and 66-73% with RSV and PVM. The region between aa residues 160 and 340 appears to be highly conserved among metapneumoviruses and to a somewhat lesser extent the Pneumovirinae (Miyahara et al., 1992; Li et al., 1996; Barr et al., 1991). This is in agreement with MPV being a metapneumovirus, showing 100% similarity with APV C.

# The phosphoprotein (P) gene

The second ORF in the genome map codes for a 294 aa protein which shares 68% as sequence homology with the P protein of APV-C, and only 22-26% with the P protein of RSV (Table 6). The P gene of MPV contains one substantial ORF and in that respect is similar to P from many other paramyxoviruses (Reviewed in Lamb and Kolakofsky, 1996; Sedlmeier et al., 1998). In contrast to APV A and B and PVM and similar to RSV and APV-C the MPV P ORF lacks cysteine residues. Ling (1995) suggested that a region of high similarity between all pneumoviruses (aa 185-241) plays a role in either the RNA synthesis process or in maintaining the structural integrity of the nucleocapsid complex. This region of high similarity is also found in MPV (Figure 9) especially when conservative substitutions are taken in account, showing 100% similarity 20 with APV-C, 93 % with APV-A and B, and approximately 81% with RSV. The Cterminus of the MPV P protein is rich in glutamate residues as has been ---------

#### The matrix (M) protein gene

described for APVs (Ling et al., 1995).

25 The third ORF of the MPV genome encodes a 254 aa protein, which resembles the M ORFs of other pneumoviruses. The M ORF of MPV has exactly the same size as the M ORFs of other metapneumoviruses (Table 5) and shows high aa sequence homology with the matrix proteins of APV (78-87%), lower homology with those of RSV and PVM (37-38%) and 10% or less homology with those of 30 other paramyxoviruses (Table 6). Easton (1997) compared the sequences of matrix proteins of all pneumoviruses

and found a conserved heptadpeptide at residue 14 to 19 that is also conserved in MPV (Figure 10). For RSV, PVM and APV small secondary ORFs within or overlapping with the major ORF of M have been identified (52 aa and 51 aa in bRSV, 75 aa in RSV, 46 aa in PVM and 51 aa in APV) (Yu et al., 1992; Easton et

al., 1997; Samal et al., 1991; Satake et al., 1984). We noticed two small ORFs in the M ORF of MPV. One small ORF of 54 aa residues was found within the major M ORF (fragment 1, Figure 7), starting at nucleotide 2281 and one small ORF of 33 aa residues was found overlapping with the major ORF of M starting at nucleotide 2893 (fragment 2, Figure 7). Similar to the secondary ORFs of RSV and APV there is no significant homology between these secondary ORFs and secondary ORFs of the other pneumoviruses, and apparent start or stop signals are lacking. In addition, evidence for the synthesis of proteins corresponding to these secondary ORFs of APV and RSV has not been reported.

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## The fusion protein (F) gene

The F ORF of MPV is located adjacent to the M ORF, which is characteristic for members of the *Metapneumovirus* genus. The F gene of MPV encodes a 539 aa protein, which is two aa residues longer than F of APV-C (Table 5). Analysis of the aa sequence revealed 81% homology with APV-C, 67% with APV-A and B, 33-39% with pneumovirus F proteins and only 10-18% with other paramyxoviruses (Table 6). One of the conserved features among F proteins of paramyxoviruses, and also seen in MPV is the distribution of cysteine residues (Morrison, 1988; Yu et al., 1991). The metapneumoviruses share 12 cysteine residues in F1 (7 are conserved among all paramyxoviruses), and two in F2 (1 is conserved among all paramyxoviruses). Of the 3 potential N-linked glycosylation sites present in the F ORF of MPV, none are shared with RSV and two (position 74 and 389) are shared with APV. The third, unique, potential N-linked glycosylation site for MPV is located at position 206 (Figure 11).

Despite the low sequence homology with other paramyxoviruses, the F protein of MPV revealed typical fusion protein characteristics consistent with those described for the F proteins of other Paramyxoviridae family members (Morrison, 1988). F proteins of Paramyxoviridae members are synthesized as inactive precursors (F0) that are cleaved by host cell proteases which generate amino terminal F2 subunits and large carboxy terminal F1 subunits. The proposed cleavage site (Collins et al., 1996) is conserved among all members of the Paramyxoviridae family. The cleavage site of MPV contains the residues RQSR. Both arginine (R) residues are shared with APV and RSV, but the glutamine (Q) and serine (S) residues are shared with other paramyxoviruses

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such as human parainfluenza virus type 1, Sendai virus and morbilliviruses (data not shown).

The hydrophobic region at the amino terminus of F1 is thought to function as the membrane fusion domain and shows high sequence similarity among paramyxoviruses and morbilliviruses and to a lesser extend the pneumoviruses (Morrison, 1988). These 26 residues (position 137-163, Figure 11) are conserved between MPV and APV-C, which is in agreement with this region being highly conserved among the metapneumoviruses (Naylor et al., 1998; Seal et al., 2000). As is seen for the F2 subunits of APV and other paramyxoviruses, MPV revealed a deletion of 22 aa residues compared with RSV (position 107-128, Figure 11). Furthermore, for RSV and APV, the signal peptide and anchor domain were found to be conserved within subtypes and displayed high variability between subtypes (Plows et al., 1995; Naylor et al., 1998). The signal peptide of MPV (aa 10-35, Figure 11) at the amino terminus of F2 exhibits some sequence similarity with APV-C (18 out of 26 as residues are similar) and less conservation with other APVs or RSV. Much more variability is seen in the membrane anchor domain at the carboxy terminus of F1, although some homology is still seen with APV-C.

#### 20 The 22K (M2) protein

The M2 gene is unique to the *Pneumovirinae* and two overlapping ORFs have been observed in all pneumoviruses. The first major ORF represents the M2-1 protein which enhances the processivity of the viral polymerase (Collins *et al.*, 1995; Collins, 1996) and its readthrough of intergenic regions (Hardy *et al.*, 1998; Fearns *et al.*, 1999). The M2-1 gene for MPV, located adjacent to the F gene, encodes a 187 aa protein (Table 5), and reveals the highest (84%) homology with M2-1 of APV-C (Table 6). Comparison of all pneumovirus M2-1 proteins revealed the highest conservation in the amino-terminal half of the protein (Collins *et al.*, 1990; Zamora *et al.*, 1992; Ahmadian *et al.*, 1999), which is in agreement with the observation that MPV displays 100% similarity with APV-C in the first 80 aa residues of the protein (Figure 12A). The MPV M2-1 protein contains 3 cysteine residues located within the first 30 aa residues that are conserved among all pneumoviruses. Such a concentration of cysteines is frequently found in zinc-binding proteins (Ahmadian *et al.*, 1991; Cuesta et al., 2000).

The secondary ORFs (M2-2) that overlap with the M2-1 ORFs of pneumoviruses are conserved in location but not in sequence and are thought to be involved in the control of the switch between virus RNA replication and transcription (Collins et al., 1985; Elango et al., 1985; Baybutt et al., 1987; Collins et al., 1990; Ling et al., 1992; Zamora et al., 1992; Alansari et al., 1994; Ahmadian et al., 1999; Bermingham et al., 1999). For MPV, the M2-2 ORF starts at nucleotide 512 in the M2-1 ORF (Figure 7), which is exactly the same start position as for APV-C. The length of the M2-2 ORFs are the same for APV-C and MPV, 71 aa residues (Table 5). Sequence comparison of the M2-2 ORF (Figure 12B) revealed 64% aa sequence homology between MPV and APV-C and only 44-48% aa sequence homology between MPV and APV-A and B (Table 6).

## The small hydrophobic (SH) protein gene

The gene located adjacent to M2 of MPV encodes a 183 aa protein (Figure 7). 15 Analysis of the nucleotide sequence and its deduced amino acid sequence revealed no discernible homology with other RNA virus genes or gene products. The SH ORF of MPV is the longest SH ORF known to date (Table 5). The composition of the aa residues of the SH ORF is relatively similar to that of APV, RSV and PVM, with a high percentage of threonine and serine residues which may represent O-linked glycosylation sites (22%, 18%, 19%, 20.0%, 21% and 20 28% serine/threonine content for MPV, APV, RSV A, RSV B, bRSV and PVM respectively). The SH ORF of MPV contains 10 cysteine residues, whereas APV SH contains 16 cysteine residues. All pneumoviruses have similar numbers of potential N-glycosylation sites (MPV 2, APV 1, RSV 2, bRSV 3, PVM 4). The hydrophobicity profiles for the MPV SH protein and SH of APV and RSV 25 revealed similar structural characteristics (Figure 13B). The SH ORFs of APV and MPV have a hydrophylic N-terminus (aa 1-30), a central hydrophobic domain (aa 36-53) which can serve as a potential membrane spanning domain, a second hydrophobic domain around residue 160 and a hydrophilic C-terminus. In contrast, RSV SH appears to lack the C-terminal half of the APV and MPV 30 ORFs. In all pneumovirus SH proteins the hydrophobic domain is flanked by basic amino acids, which are also found in the SH ORF for MPV (aa 35 and 54).

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# The attachment glycoprotein (G)

The G ORF of MPV is located adjacent to the SH gene and encodes a 236 amino acid protein. A secondary small ORF is found immediately following this ORF, potentially coding for 68 aa residues (pos. 6973-7179, fragment 3, Figure 7), but lacking a start codon. A third major ORF, in a different reading frame, of 194 aa 5 residues (fragment 4, Figure 7) is overlapping with both of these ORFs, starting with an ATG codon at nucleotide 6416. This major ORF is followed by a fourth ORF in the same reading frame, possibly coding for 65 aa residues (pos. 7002-7198, fragment 5, Figure 7) but again lacking a start codon. Unlike the first ORF, the other ORFs do not have apparent gene start or gene end sequences (see 10 below). Although the 236 aa residue G ORF probably represents at least a part of the MPV attachment protein it can not be excluded that the additional coding sequences are expressed as separate proteins or as part of the attachment protein through some RNA editing event. It should be noted that for APV and 15 RSV no secondary ORFs after the primary G ORF have been identified but that both APV and RSV have secondary ORFs within the major ORF of G. However, evidence for expression of these ORFs is lacking and there is no homology between the predicted as sequences for different viruses (Ling et al., 1992). The secondary ORFs in MPV G do not reveal characteristics of other G proteins and whether the additional ORFs are expressed requires further investigation. 20 BLAST analyses with all four ORFs revealed no discernible homology at the nucleotide or as sequence level with other known virus genes or gene products. This is in agreement with the low sequence homologies found for other G proteins such as hRSV A and B (53%) (Johnson et al., 1987) and APV A and B 25 (38%) (Juhasz et al., 1994). Whereas most of the MPV ORFs resemble those of APV both in length and sequence, the GORF of MPV is considerably smaller than the GORF of APV (Table 5). The aa sequence revealed a serine and threonine content of 34%, which is even higher than the 32% for RSV and 24% for APV. The G ORF also contains 8.5% proline residues, which is higher than the 8% for RSV and 7% for APV. The 30 unusual abundance of proline residues in the G proteins of APV, RSV and MPV has also been observed in glycoproteins of mucinous origin where it is a major

determinant of the proteins three dimensional structure (Collins et al., 1983;

Wertz et al., 1985; Jentoft, 1990).

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The number of potential *N*-linked glycosylation sites in G of MPV is similar to other pneumoviruses: MPV has 5, whereas hRSV has 7, bRSV has 5, and APV has 3 to 5.

The predicted hydrophobicity profile of MPV G revealed characteristics similar to the other pneumoviruses. The amino-terminus contains a hydrophylic region followed by a short hydrophobic area (aa 33-53) and a mainly hydrophilic carboxy terminus (Figure 14B). This overall organisation is consistent with that of an anchored type II transmembrane protein and corresponds well with these regions in the G protein of APV and RSV. The G ORF of MPV contains only 1 cysteine residue in contrast to RSV and APV (5 and 20 respectively).

### The polymerase gene (L)

In analogy to other negative strand viruses, the last ORF of the MPV genome is the RNA-dependent RNA polymerase component of the replication and transcription complexes. The L gene of MPV encodes a 2005 as protein, which is 1 residue longer than the APV-A protein (Table 5). The L protein of MPV shares 64% homology with APV-A, 42-44% with RSV, and approximately 13% with other paramyxoviruses (Table 6). Poch et al. (1989; 1990) identified six conserved domains within the L proteins of non-segmented negative strand RNA viruses, from which domain III contained the four core polymerase motifs that are thought to be essential for polymerase function. These motifs (A, B, C and D) are well conserved in the MPV L protein: in motifs A, B and C: MPV shares 100% similarity with all pneumoviruses and in motif D MPV shares 100 % similarity with APV and 92% with RSVs. For the entire domain III (aa 627-903 in the L ORF), MPV shares 77% identity with APV, 61-62% with RSV and 23-27% with other paramyxoviruses (Figure 15). In addition to the polymerase motifs the pneumovirus L proteins contain a sequence which conforms to a consensus ATP binding motif K(X)21GEGAGN(X)20K (Stec, 1991). The MPV L ORF contains a similar motif as APV, in which the spacing of the intermediate residues is off by one: K(x)22GEGAGN(X)19 K.

## Phylogenetic analyses

As an indicator for the relationship between MPV and members of the *Pneumovirinae*, phylogenetic trees based on the N, P, M, and F ORFs have been constructed previously (van den Hoogen *et al.*, 2001) and revealed a close

relationship between MPV and APV-C. Because of the low homology of the MPV SH and G genes with those of other paramyxoviruses, reliable phylogenetic trees for these genes can not be constructed. In addition, the distinct genomic organization between members of the Pneumovirus and Metapneumovirus genera make it impossible to generate phylogenetic trees based on the entire genomic sequence. We therefore only constructed phylogenetic trees for the M2 and L genes in addition to those previously published. Both these trees confirmed the close relation between APV and MPV within the Pneumovirinae subfamily (Figure 16).

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### MPV non-coding sequences

The gene junctions of the genomes of paramyxoviruses contain short and highly conserved nucleotide sequences at the beginning and end of each gene (gene start and gene end signals), possibly playing a role in initiation and termination of transcription (Curran et al., 1999). Comparing the intergenic sequences between all genes of MPV revealed a consensus sequence for the gene start signal of the N. P. M. F. M2 and G: GGGACAAGU (Figure 17A), which is identical to the consensus gene start signal of the metapneumoviruses (Ling et al., 1992; Yu et al., 1992; Li et al., 1996; Bäyon-Auboyer et al., 2000). The gene start signals for the SH and L genes of MPV were found to be slightly different from this consensus (SH: GGGAUAAAU, L: GAGACAAAU). For APV the gene start signal of L was also found to be different from the consensus: AGGACCAAT (APV-A) (Randhawa et al., 1996) and GGGACCAGT (APV-D) (Bäyon-Auboyer et al., 2000).

In contrast to the similar gene start sequences of MPV and APV, the consensus gene end sequence of APV, UAGUUAAUU (Randhawa et al., 1996), could not be found in the MPV intergenic sequences. The repeated sequence found in most genes, except the G-L intergenic region, was U AAAAA U/A/C, which could possibly act as gene end signal. However, since we sequenced viral RNA rather than mRNA, definitive gene end signals could not be assigned and thus requires further investigation. The intergenic regions of pneumoviruses vary in size and sequence (Curran et al., 1999; Blumberg et al., 1991; Collins et al., 1983;). The intergenic regions of MPV did not reveal homology with those of APV and RSV and range in size from 10 to 228 nucleotides (Figure 17B). The intergenic region

between the M and F ORFs of MPV contains part of a secondary ORF, which 35

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starts in the primary M ORF (see above). The intergenic region between SH and G contains 192 nucleotides, and does not appear to have coding potential based on the presence of numerous stop-codons in all three reading frames. The intergenic region between G and L contains 241 nucleotides, which may include additional ORFs (see above). Interestingly, the start of the LORF is located in these secondary ORFs. Whereas the L gene of APV does not start in the preceding G ORF, the L ORF of RSV also starts in the preceding M2 gene. At the 3' and 5'extremities of the genome of paramyxoviruses short extragenic region are referred to as the leader and trailer sequences, and approximately the first 12 nucleotides of the leader and last 12 nucleotides of the trailer are complementary, probably because they each contain basic elements of the viral promoter (Curran et al., 1999; Blumberg et al., 1991; Mink et al., 1986). The 3'leader of MPV and APV are both 41 nucleotides in length, and some homology is seen in the region between nucleotide 16 and 41 of both viruses (18 out of 26 nucleotides) (Figure 17B). As mentioned before the first 15 nucleotides of the MPV genomic map are based on a primer sequence based on the APV genome. The length of the 5'trailer of MPV (188 nucleotides) resembles the size of the RSV 5'trailer (155 nucleotides), which is considerably longer than that of APV (40 nucleotides). Alignments of the extreme 40 nucleotides of the trailer of MPV and the trailer of APV revealed 21 out of 32 nucleotides homology, apart from the extreme 12 nucleotides which represent primer sequences based on the genomic sequence of APV. Our sequence analyses revealed the absence of NS1 and NS2 genes at the 3'end of the genome and a genomic organisation resembling the organisation of metapneumoviruses (3'-N-P-M-F-M2-SH-G-L-5'). The high sequence homology found between MPV and APV genes further emphasises the close relationship between these two viruses. For the N, P, M, F, M2-1 and M2-2 genes of MPV an overall amino acid homology of 79% is found with APV-C. In fact, for these genes APV-C and MPV revealed sequence homologies which are in the same range as sequence homologies found between subgroups of other genera, such as RSV- A and B or APV-A and B. This close relationship between APV-C and MPV is also seen in the phylogenetic analyses which revealed MPV and APV-C always in the same branch, separate from the branch containing APV-A and B. The identical genomic organisation, the sequence homologies and phylogentic analyses are all in favour of the classification of MPV as the first member in the Metapneumovirus genus that is

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isolatable from mammals. It should be noted that the found sequence variation between different virus isolates of MPV in the N, M, F and L genes revealed the possible existence of different genotypes (van den Hoogen et al., 2001). The close relationship between MPV and APV-C is not reflected in the host range, since APV infects birds in contrast to MPV (van den Hoogen et al., 2001). This difference in host range may be determined by the differences between the SH and G proteins of both viruses that are highly divergent. The SH and G proteins of MPV did not reveal significant as sequence homology with SH and G proteins of any other virus. Although the amino acid content and hydrophobicity plots are in favour of defining these ORFs as SH and G, experimental data are required to assess their function. Such analyses will also shed light on the role of the additional overlapping ORFs in these SH and G genes. In addition, sequence analyses on the SH and G genes of APV-C might provide more insight in the function of the SH and G proteins of MPV and their relationship with those of APV-C. The noncoding regions of MPV were found to be fairly similar to those of APV. The 3'leader and 5' trailer sequences of APV and MPV displayed a high degree of homology. Although the lengths of the intergenic regions were not always the same for APV and MPV, the consensus gene start signals of most of the ORFs were found to be identical. In contrast, the gene end signals of APV were not found in the MPV genome. Although we did find a repetitive sequence (U AAAAA U/A/C) in most intergenic regions, sequence analysis of viral mRNAs. is required to formally delineate those gene end sequences. It should be noted that sequence information for 15 nucleotides at the extreme 3'end and 12 nucleotides at the extreme 5'end is obtained by using modified rapid amplification of cDNA ends (RACE) procedures. This technique has been proven to be successful by others for related viruses (Randhawa, J.S. et al., Rescue of synthetic minireplicons establishes the absence of the NS1 and NS2 genes from avian pneumovirus. J. Virol, 71, 9849-9854 (1997); Mink, M.A., et al. Nucleotide sequences of the 3' leader and 5' trailer regions of human respiratory syncytial virus genomic RNA. Virology 185, 615-24 (1991).) To determine the sequence of the 3' vRNA leader sequence, a homopolymer A tail is added to purified vRNA using poly-A-polymerase and the leader sequence subsequently amplified by PCR using a poly-T primer and a primer in the N gene. To determine the sequence of the 5' vRNA trailer sequence, a cDNA copy of the trailer sequence is made using reverse transcriptase and a primer in the L gene, followed by

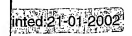
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homopolymer dG tailing of the cDNA with terminal transferase. Subsequently, the trailer region is amplified using a poly-C primer and a primer in the L gene. As an alternative strategy, vRNA is ligated to itself or synthetic linkers, after which the leader and trailer regions are amplified using primers in the L and N genes and linker-specific primers. For the 5' trailer sequence direct dideoxynucleotide sequencing of purified vRNA is also feasible (Randhawa, 1997). Using these approaches, we can analyse the exact sequence of the ends of the hMPV genome. The sequence information provided here is of importance for the generation of diagnostic tests, vaccines and antivirals for MPV and MPV infections.

### Materials and Methods

Sequence analysis

Virus isolate 00-1 was propagated to high titers (approximately 10,000 15 TCID50/ml) on tertiary monkey kidney cells as described previously (van den Hoogen et al., 2001). Viral RNA was isolated from supernatants from infected cells using a High Pure RNA Isolating Kit according to instructions from the manufacturer (Roch Diagnostics, Almere, The Netherlands). Primers were 20 designed based on sequences published previously (van den Hoogen et al., 2001) in addition to sequences published for the leader and trailer of APV/RSV (Randhawa et al., 1997; Mink et al., 1991) and are available upon request. RT-PCR assays were conducted with viral RNA, using a one-tube assay in a total volume of 50µl with 50 mM Tris pH 8.5, 50 mM NaCl, 4.5 mM MgCl<sub>2</sub>, 2 mM DTT. 1 µM forward primer, 1µM reverse primer, 0.6 mM dNTP's, 20 units 25 RNAsin (Promega, Leiden, The Netherlands), 10 U AMV reverse transcriptase (Promega, Leiden, The Netherlands), and 5 units Taq Polymerase (PE Applied Biosystems, Nieuwerkerk aan de IJssel, The Netherlands). Reverse transcription was conducted at 42°C for 30 minutes, followed by 8 minutes inactivation at 95°C. The cDNA was amplified during 40 cycles of 95°C, 1 min.; 42°C, 2 30 min.72°C, 3 min. with a final extension at 72°C for 10 minutes. After examination on a 1% agarose gel, the RT-PCR products were purified from the gel using a Qiaquick Gel Extraction kit (Qiagen, Leusden, The Netherlands) and sequenced directly using a Dyenamic ET terminator sequencing kit (Amersham Pharmacia Biotech, Roosendaal, the Netherlands) and an ABI 373 automatic 35



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DNA sequencer (PE Applied Biosystem, Nieuwerkerk aan den IJssel, the Netherlands), according to the instructions of the manufacturer. Sequence alignments were made using the clustal software package available in the software package of BioEdit version5.0.6. (http://jwbrown.mbio.ncsu.edu/Bioedit//bioedit.html; Hall, 1999).

### Phylogenetic analysis

To construct phylogenetic trees, DNA sequences were aligned using the ClustalW software package and maximum likelihood trees were generated using the DNA-ML software package of the Phylip 3.5 program using 100 bootstraps and 3 jumbles. Bootstrap values were computed for consensus trees created with the consense package (Felsenstein, 1989).

The MPV genomic sequence is available from Genbank under accession number AF371337. All other sequences used here are available from Genbank under accession numbers AB046218 (measles virus, all ORFs), NC-001796 (human parainfluenza virus type 3, all ORFs), NC-001552 (Sendai virus, all ORFs), X57559 (human parainfluenza virus type 2, all ORFs), NC-002617 (New Castle

- Disease virus, all ORFs), NC-002728 (Nipah virus, all ORFs), NC-001989 (bRSV, all ORFs), M11486 (hRSV A, all ORFs except L), NC-001803 (hRSV, L ORF), NC-001781 (hRSV B, all ORFs), D10331 (PVM, N ORF), U09649 (PVM, P ORF), U66893 (PVM, M ORF), U66893 (PVM, SH ORF), D11130 (PVM, G ORF), D11128 (F ORF). The PVM M2 ORF was taken from Ahmadian (1999), AF176590 (APV-C, N ORF), U39295 (APV-A, N ORF), U39296 (APV-B, N ORF), AF262571 (APV-C, M ORF), U37586 (APV-B, M ORF), X58639 (APV-A, M ORF),
- AF176591 (APV-C, P ORF), AF325443 (APV-B, P ORF), U22110 (APV-A, P ORF), AF187152 (APV-C, F ORF), Y14292 (APV-B, F ORF), D00850 (APV-A, F ORF), AF176592 (APV-C, M2 ORF), AF35650 (APV-B, M2 ORF), X63408 (APV-A, M2 ORF), U65312 (APV-A, L ORF), S40185 (APV-A, SH ORF).



Table 5: Lengths of the ORFs of MPV and other paramyxoviruses.

	$N^{i}$	P	M	F	M2-	M2-	SH	G	$\mathbf{L}$
					1	2			
MPV	394	294	254	539	187	71	183	236	2005
APV A	391	278	254	538	186	73	174	391	2004
APV B	391	279	254	538	186	<b>7</b> 3	_2	414	_2
APV C	394	294	254	537	184	71	_2	_2	_2
APV D	_2	_2	_2	_2	_2	_2	_2	389	_2
hRSV A	391	241	256	574	194	90	64	298	2165
hRSV B	391	241	249	574	195	93	65	299	2166
bRSV	391	241	256	569	186	93	81	257	2162
PVM	393	295	257	537	176	77	92	396	_2
others <sup>3</sup>	418-	225-	335-	539-	_4	_4	_4	-4	2183-
	<b>542</b>	709	393	565					2262

# Footnotes:

- 1. length in amino acid residues.
- 2. sequences not available
- others: human parainfluenza virus type 2 and 3, Sendai virus, measles virus,
   nipah virus, phocine distemper virus, and New Castle Disease virus.
  - 4. . ORF not present in viral genome

Table 6: Amino acid sequence identity between the ORFs of MPV and those of other paramyxoviruses<sup>1</sup>.

	N	P	M	F	M2-	M2-	L
					1	2	
APV A	69	55	78	67	72	48	64
APV B	69	51	76	67	71	44	_2
APV C	88	68	87	81	84	64	_2
hRSV A	42	24	38	34	36	32	42
hRSV B	41	23	37	33	35	32	44
bRSV	42	22	38	34	35	30	44
PVM	45	26	<b>37</b>	39	33	30	<b>_2</b> .
others <sup>3</sup>	7-11	4-9	7-10	10-	4	_4	13-
				18			14

# Footnotes:

- 1. No sequence homologies were found with known G and SH proteins and were
- 10 thus excluded
  - 2. Sequences not available.
  - 3. See list in table 5, footnote 3.
  - 4. ORF absent in viral genome.

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Primers used for RT-PCR detection of known paramyxo-viruses. Primers for hPIV-1 to 4, mumps, measles, Tupaia, Mapuera and Hendra are developed in house and based on allignments of available sequences. Primers for New Castle Disease Virus are taken from Seal, J., J. et al; Clin. Microb., 2624-2630, 1995.

5 Primers for Nipah and general paramyxovirus-PCR are taken from: Chua, K.B., et al; Science, 288 26 may 2000

	Virus		primers	located	l in protein
	HPIV-1	fwd	5'-TGTTGTCGAGACTATTCCAA-3'		HN
10		Rev	5'-TGTTG(T/A)ACCAGTTGCAGTCT-3'		
	HPIV-2	Fwd	5'-TGCTGCTTCTATTGAGAAACGCC-3'	N	
		Rev	5'-GGTGAC/T TC(T/C)AATAGGGCCA-3'		
	HPIV-3	$\mathbf{Fwd}$	5'-CTCGAGGTTGTCAGGATATAG-3'		HN
		Rev	5'-CTTTGGGAGTTGAACACAGTT-3'		
15	HPIV-4	Fwd	5'-TTC(A/G)GTTTTAGCTGCTTACG-3'	N	
		Rev	5'-AGGCAAATCTCTGGATAATGC-3'		
	Mumps	Fwd	5'-TCGTAACGTCTCGTGACC-3'	SH	
		Rev	5'-GGAGATCTTTCTAGAGTGAG-3'		
	NDV	Fwd	5'-CCTTGGTGA:TCTATCCGIAG-3'		F
20		Rev	5'-CTGCCACTGCTAGTTGiGATAATCC-3		
	Tupaia	Fwd	5'-GGGCTTCTAAGCGACCCAGATCTTG-	.3'	N
		Rev	5'-GAATTTCCTTATGGACAAGCTCTGTG	fC-3,	
	Mapuera	Fwd	5'-GGAGCAGGAACTCCAAGACCTGGAG	-3 <b>'</b>	N
		Rev:	5'-GCTCAACCTCATCACATACTAACCC-S	3'	
25	Hendra	Fwd	5'-GAGATGGGCGGCAAGTGCGGCAAC	JAG-3'	N
		$\mathbf{Rev}$	5'-GCCTTTGCAATCAGGATCCAAATTTG	;GG-3°	
	Nipah	Fwd	5'-CTGCTGCAGTTCAGGAAACATCAG-3	3'	N
		Rev	5'-ACCGGATGTGCTCACAGAACTG-3'		
	HRSV	$\mathbf{Fwd}$	5'-TTTGTTATAGGCATATCATTG-3'		F
30		Rev	5'-TTAACCAGCAAAGTGTTA-3'		
	Measles	Fwd	5'-TTAGGGCAAGAGATGGTAAGG-3'		N
		$\mathbf{Rev}$	5'-TTATAACAATGATGGAGGG-3'		
	General Par	amyxov	riridae :		
		Fwd	5'-CATTAAAAAGGGCACAGACGC-3'		P
35		Rev	5'-TGGACATTCTCCGCAGT-3'		

#### Primers for RAP-PCR:

ZF1: 5'-CCCACCACCAGAGAGAAA-3'

ZF4: 5'-ACCACCAGAGAGAAACCC-3'

5 ZF7: 5'-ACCAGAGAGAAACCCACC-3'

ZF10: 5'-AGAGAGAAACCCACCACC-3'

ZF13: 5'-GAGAAACCCACCACCAGA-3'

ZF16: 5'-AAACCCACCACCAGAGAG-3'

10 CS1: 5'-GGAGGCAAGCGAACGCAA-3'

CS4: 5'-GGCAAGCGAACGCAAGGA-3'

CS7: 5'-AAGCGAACGCAAGGAGGC-3'

CS10:5'-CGAACGCAAGGAGGCAAG-3'

CS13:5'-ACGCAAGGAGGCAAGCGA-3'

15 CS16:5'-CAAGGAGGCAAGCGAACG-3'

20 fragments successfully purified and sequenced:

#### 10 fragments found with sequence homology in APV

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	Fragment 1	ZF 7, 335 bp	N gene
	Fragment 2	ZF 10, 235 bp	N gene
	Fragment 3	ZF 10, 800 bp	M gene
	Fragment 4	CS 1, 1250 bp	F gene
25	Fragment 5	CS 10, 400 bp	F gene
	Fragment 6	CS 13, 1450 bp	F gene
	Fragment 7	CS 13, 750 bp	F gene
	Fragment 8	ZF 4, 780 bp	L gene (protein level)
	Fragment 9	ZF 10, 330 bp	L gene (protein level)
30	Fragment 10	ZF10, 250bp	L gene (protein level)

Primers used for RAP-PCR amplification of nucleic acids from the prototype isolate.

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#### Addendum

#### Background information on Pneumovirinae

The family of Paramyxoviridae contains two subfamilies: the Paramyxoviriae and the Pneumoviriae. The subfamily Pneumoviriae consists of two genera: Pneumovirus and Metapneumovirus. The genus Pneumovirus contains the human, bovine, ovine and caprine respiratory syncytial viruses and the pneumonia virus of mice (PVM). The genus Metapneumovirus contains the avian pneumoviruses (APV, also referred to as TRTV).

The classification of the genera in the subfamily *Pneumovirinae* is based on classical virus characteristics, gene order and gene constellation. Viruses of the genus *Pneumovirus* are unique in the family of *Paramyxoviridae* in having two nonstructural proteins at the 3'end of the genome (3'-NS1-NS2-N-P-M-SH-G-F-M2-L-5'). In contrast, viruses in the genus *Metapneumovirus* lack the NS1 and NS2 genes and the organisation of genes between the M and L coding regions is different: 3'-N-P-M-F-M2-SH-G-L-5'.

All members of the subfamily Paramyxovirinae have haemagluttinating activity, but this function is not a defining feature for the subfamily Pneumovirinae, being absent in RSV and APV but present in PMV. Neuraminidase activity is present in members of the genera Paramyxovirus and Rubulavirus (subfamily Paramyxovirinae) but is absent in the genus Morbillivirus (subfamily Paramyxovirinae) and the genera Pneumovirus and Metapneumovirus (subfamily Pneumovirinae).

A second distinguishing feature of the subfamily *Pneumovirinae* is the apparent limited utilization of alternative ORFs within mRNA by RSV. In contrast, several members of the subfamily *Paramyxovirinae*, such as Sendai and Measles viruses, access alternative ORFs within the mRNA encoding the phosphoprotein

(P) to direct the synthesis of a novel protein.

The G protein of the *Pneumovirinae* does not have sequence relatedness or structural similarity to the HN or H proteins of *Paramyxovirinae* and is only approximately half the size of their chain length. In addition, the N and P proteins are smaller than their counterparts in the *Paramyxovirinae* and lack

35 unambigous sequence homology. Most nonsegmented negative stranded RNA

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viruses have a single matrix (M) protein. Members of the subfamily *Pneumovirinae* are an exception in having two such proteins, M and M2. The M protein is smaller than its *Paramyxovirinae* counterparts and lacks sequence relatedness with *Paramyxovirinae*.

When grown in cell cultures, members of the subfamily *Pneumovirinae* show typical cytopathic effects; they induce characteristic syncytia formation of cells. (Collins, 1996).

#### 10 The subfamily Pneumovirinae, genus Pneumovirus

hRSV is the type-species of the genus *Pneumovirus* and is a major and widespread cause of lower respiratory tract illness during infancy and early childhood (Selwyn, 1990). In addition, hRSV is increasingly recognised as an important pathogen in other patient groups, including immune compromised individuals and the elderly. RSV is also an important cause of community-acquired pneumonia among hospitalised adults of all ages (Englund, 1991; Falsey, 2000; Dowell, 1996).

Two major antigenic types for RSV (A and B) have been identified based on differences in their reactivity with monoclonal and polyclonal antibodies and by nucleic acid sequence analyses (Anderson, 1985; Johnson, 1987; Sullender, 2000). In particular the G protein is used in distinguishing the two subtypes. RSV-A and B share only 53% amino acid sequence homology in G, whereas the other proteins show higher homologies between the subtypes (table 1) (Collins, 1996). Detection of RSV infections has been described using monoclonal and polyclonal antibodies in immunofluorescence techniques (DIF, IFA), virus neutralisation assays and ELISA or RT-PCR assays (Rothbarth, 1988; Van Milaan, 1994; Coggins, 1998).

Closely related to hRSV are the bovine (bRSV), ovine (oRSV) and caprine RSV (oRSV), from which bRSV has been studied most extensively. Based on sequence homology with hRSV, the ruminant RSVs are classified within the *Pneumovirus* genus, subfamily *Pneumovirinae* (Collins, 1996). Diagnosis of ruminant RSV infection and subtyping is based on the combined use of serology, antigen detection, virus isolation and RT-PCR assays (Uttenthal, 1996; Valarcher, 1999; Oberst, 1993; Vilcek, 1994).

Several analyses on the molecular organisation of bRSV have been performed using human and bovine antisera, monoclonal antibodies and cDNA probes. These analyses revealed that the protein composition of hRSV and bRSV are very similar and the genomic organisation of bRSV resembles that of hRSV. For both bRSV and hRSV, the G and F proteins represent the major neutralisation and protective antigens. The G protein is highly variable between the hRSV subtypes and between hRSV and bRSV (53 and 28% respectively) (Prozzi, 1997; Lerch, 1990). The F proteins of hRSV and bRSV strains present comparable structural characteristics and antigenic relatedness. The F protein of bRSV shows 80-81% homology with hRSV, while the two hRSV subtypes share 90% homology in F (Walravens, K. 1990).

Studies based on the use of hRSV and bRSV specific monoclonal antibodies have suggested the existence of different antigenic subtypes of bRSV. Subtypes A, B, and AB are distinguished based on reaction patterns of monoclonal antibodies

and AB are distinguished based on reaction patterns of monoclonal antibodies

specific for the G protein (Furze, 1994; Prozzi, 1997; Elvander, 1998). The
epidemiology of bRSV is very similar to that of hRSV. Spontaneous infection in
young cattle is frequently associated with severe respiratory signs, whereas
experimental infection generally results in milder disease with slight pathologic
changes (Elvander, 1996).

RSV has also been isolated from naturally infected sheep (oRSV) (LeaMaster, 1983) and goats (cRSV) (Lehmkuhl, 1980). Both strains share 96% nucleotide sequence with the bovine RSV and are antigenically crossreacting. Therefore, these viruses are also classified within the *Pneumovirus* genus.

A distinct member of the subfamily *Pneumovirinae*, genus *Pneumovirus* is the Pneumonia virus of mice (PVM).

PVM is a common pathogen in laboratory animal colonies, particularly those containing atymic mice. The naturally acquired infection is thought to be asymptomatic, though passage of virus in mouse lungs resulted in overt signs of disease ranging from an upper respiratory tract infection to a fatal pneumonia (Richter, 1988; Weir, 1988).

Restricted serological crossreactivity between the nucleocapsid protein (N) and the phosphoprotein (P) of PVM and hRSV has been described but none of the external proteins show cross-reactivity, and the viruses can be distinguished from each other in virus neutralisation assays (Chambers, 1990a; Gimenez,

35 1984; Ling, 1989a).

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The glycoproteins of PVM appear to differ from those of other paramyxoviruses and resemble those of RSV in terms of their pattern of glycosylation. They differ, however, in terms of processing. Unlike RSV, but similar to the other paramyxoviruses, PVM has haemagglutinating activity with murine erythrocytes, for which the G protein appears to be responsible since a monoclonal antibody to this protein inhibits haemagglutination (Ling, 1989b). The genome of PVM resembles that of hRSV, including two nonstructural proteins at its 3'end and a similar genomic organisation (Chambers, 1990a; Chambers, 1990b). The nucleotide sequences of the PVM NS1/NS2 genes are not detectably homologous with those of hRSV (Chambers, 1991). Some proteins of PVM show strong homology with hRSV (N: 60%, and F: 38 to 40%) while G is distinctly different (the amino acid sequence is 31 % longer) (Barr, 1991; Barr, 1994; Chambers, 1992). The PVM P gene, but not that of RSV or APV, has been reported to encode a second ORF, representing a unique PVM protein (Collins, 1996). New PVM isolates are identified by virus isolation, heamagglutination assays, virus neutralisation assay and various immuno-fluorescence techniques.

Table with addedum: Amino acid homology between the different viruses within the genus *Pneumovirus* of the subfamily *Pneumovirinae*.

Gene	hRSV's	bRSV's	oRSV v.	bRSV v.	bRSV v.	PVM vs. hRSV
			hRSV	hRSV	oRSV	
NS1	87			68-69	89	*
NS2	92			83-84	87	*
N	96		93			60
P	-		81			
M	-		89			
F	89			80-81		38-40
G	53	88-100	21-29	38-41	60-62	*
M2	92		94			41
SH	76		45-50		56	
L	-					

<sup>\*</sup> No detectable sequence homology



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#### The genus Metapneumovirus

Avian pneumoviruses (APV) has been identified as the aetiological agent of turkey rhinotracheitis (McDougall, 1986; Collins, 1988) and is therefore often referred to as turkey rhinotracheitis virus (TRTV). The disease is an upper respiratory tract infection of turkeys, resulting in high morbidity and variable, but often high, mortality. In turkey hens, the virus can also induce substantial reductions in egg production. The same virus can also infect chickens, but in this species, the role of the virus as a primary pathogen is less clearly defined, although it is commonly associated with swollen head syndrome (SHS) in

- although it is commonly associated with swollen head syndrome (SHS) in breeder chicken (Cook, 2000).
  - The virions are pleiomorphic, though mainly spherical, with sizes ranging from 70 to 600 nm and the nucleocapsid, containing the linear, non-segmented, negative-sense RNA genome, shows helical symmetry (Collins, 1986; Giraud,
- 15 1986). This morphology resembles that of members of the family Paramyxoviridae. Analyses of the APV-encoded proteins and RNAs suggested that of the two subfamilies of this family (Paramyxovirinae and Pneumovirinae), APV most closely resembled the Pneumovirinae (Collins, 1988; Ling, 1988; Cavanagh, 1988).
- APV has no non-structural proteins (NS1 and NS2) and the gene order (3'-N-P-M-F-M2-SH-G-L-5') is different from that of mammalian pneumoviruses such as RSV. APV has therefore recently been classified as the type species for the new genus *Metapneumovirus* (Pringle, 1999).
- Differences in neutralisation patterns, ELISA and reactivity with monoclonal antibodies have revealed the existence of different antigenic types of APV. Nucleotide sequencing of the G gene led to the definition of two virus subtypes (A and B), which share only 38% amino acid homology (Collins, 1993; Juhasz, 1994). An APV isolated from Colorado, USA (Cook, 1999), was shown to crossneutralize poorly with subtype A and B viruses and based on sequence
- information was designated to a novel subtype, C (Seal, 1998; Seal 2000). Two non-A/non-B APVs were isolated in France, and were shown to be antigenically distinct from subtypes A, B and C. Based on amino acid sequences of the F, L and G genes, these viruses were classified again as a novel subtype, D (Bayon-Auboyer, 2000).

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Diagnosis of APV infection can be achieved by virus isolation in chicken or turkey tracheal organ cultures (TOCs) or in Vero cell cultures. A cytopathic effect (CPE) is generally observed after one or two additional passages. This CPE is characterised by scattered focal areas of cell rounding leading to synctyial formation (Buys, 1989). A number of serology assays, including IF and virus neutralisation assays have been developed. Detection of antibodies to APV by ELISA is the most commonly used method (O'Loan, 1989; Gulati, 2000). Recently, the polymerase chain reaction (PCR) has been used to diagnose APV infections. Swabs taken from the oesophagus can be used as the starting material (Bayon-Auboyer, 1999; Shin, 2000)

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Addendum

Summary of characterisation of MPV in relation to the other paramyxoviruses.

MPV essentially does not have haemagglutinating activity.

For the genes examined, the newly discovered virus displayed less than 20% amino acid sequence with other non-pneumovirus paramyxoviruses.

Gene constellation: no NS1/NS2 at the 3'end of the genome in MPV.

Genomic organisation: different position of genes between M and L.

10 Amino acid sequence homologies found with RSV's is low: M: 37%

N: 20-22% P: 25-27% F: 32-37%

L: 30-36%

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APV is only found in birds/poultry, while we isolated our virus from mammals, in particular humans.

Absence of NS1/NS2 proteins at 3'end, just like APV

20 Absence of G protein between M and F, just like APV

Amino acid sequence homologies found with APV's is low (but higher than with

RSV):

M: 75-87%

N: 40-88%

P: 55-67%

F: 66-80%

L: 53-56%

Reason for tentative classification of the virus in the subfamily *Pneumovirinae* (in stead of *Paramyxovirinae*): sequence homology with the *Pneumovirinae*, lack of haemaglutination activity.

Reason for tentative classification of the virus in the genus *Metapneumovirinae*: higher percentage sequence homology with APV then with RSV, apparent lack of NS1 and NS2 genes at the 3' end of the viral genome and probably the same gene order as the APVs. Our PCR and sequence analyses revealed that the G gene is not located between M and F genes (similar to APV and different from RSV), but



further examination of the genomic sequence is required to determine the gene constellation and the gene order between F and L genes. It is possible that equivalents of eg NS1 and NS2 genes of MPV are found elsewhere in the viral genome.

EPO - DG 1

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118. 10, 2001

#### Claims



- An isolated essentially mammalian negative-sense single stranded RNA virus (MPV) belonging to the sub-familie Pneumovirinae of the family
   Paramyxoviridae and identifiable as phylogenetically corresponding to the genus Metapneumovirus by determining a nucleic acid sequence of said virus and testing it in phylogenetic tree analyses wherein maximum likelihood trees are generated using 100 bootstraps and 3 jumbles and finding it to be more closely phylogenetically corresponding to a virus isolate deposited as I-2614 with
   CNCM, Paris than it is corresponding to a virus isolate of avian pneumovirus (APV) also known as turkey rhinotracheitis virus (TRTV), the aetiological agent of avian rhinotracheitis..
  - A virus according to claim 1 wherein said nucleic acid sequence comprises
     an open reading frame encoding a viral protein of said virus.
    - 3. A virus according to claim 2 wherein said open reading frame is selected from the group of ORFs encoding the N, P, M, and F proteins.
- 4. A virus according to claim 2 wherein said open reading frame is selected from the group of ORFs encoding the SH or G proteins.
  - 5. A virus according to anyone of claims 1 to 4 comprising a nucleic acid or functional fragment phylogenetically corresponding to a sequence shown in figure 6a, 6b, 6c.
  - 6. A virus according to anyone of claims 1 to 5 comprising an MPV isolate deposited as I-2614 at January 19, 2001 with CNCM, Institute Pasteur, Paris or a virus isolate phylogenetically corresponding therewith.
- 7. A virus according to claim 6 isolatable from a human with respiratory tract illness.
  - 8. An isolated or recombinant nucleic acid or virus-specific functional fragment thereof obtainable from a virus according to anyone of claims 1 to 7.

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- 9. A vector comprising a nucleic acid according to claim 8.
- 10. A host cell comprising a nucleic acid according to claim 8 or a vector according to claim 9.
- 11. A proteinaceous molecule or MPV-specific functional fragment thereof encoded by a nucleic acid according to claim 8.
- 12. An antigen comprising a proteinaceous molecule or MPV-specific10 functional fragment thereof according to claim 11.
  - 13. An antibody specifically directed against an antigen according to claim 12.
- 14. A method for identifying a viral isolate as an MPV comprising reacting said viral isolate or a component thereof with an antibody according to claim 13.
  - 15. A method for identifying a viral isolate as an MPV comprising reacting said viral isolate or a component thereof with a nucleic acid according to claim 8.
- 20 16. A method according to claim 14 or 15 wherein said MPV comprises a human MPV.
  - 17. A viral isolate identifiable with a method according to anyone of claims 14 to 16 as a mammalian negative-sense single stranded RNA virus within the subfamilie *Pneumovirinae* of the family *Paramyxoviridae* and identifiable as phylogenetically corresponding to the genus *Metapneumovirus*.
  - 18. A method for virologically diagnosing an MPV infection of a mammal comprising determining in a sample of said mammal the presence of a viral isolate or component thereof by reacting said sample with a nucleic acid according to claim 8 or an antibody according to claim 13.
  - 19. A method for serologically diagnosing an MPV infection of a mammal comprising determining in a sample of said mammal the presence of an antibody specially directed against an MPV or component thereof by reacting said sample

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with a proteinaceous molecule or fragment thereof according to claim 11 or an antigen according to claim 12.

- 20. A diagnostic kit for diagnosing an MPV infection comprising a virus according to anyone of claims 1 to 7, a nucleic acid according to claim 8, a proteinaceous molecule or fragment thereof according to claim 11, an antigen according to claim 12 and/or an antibody according to claim 13.
- 21. Use of a virus according to any one claims 1 to 7, a nucleic acid according to claim 8, a vector according to claim 9, a host cell according to claim 10, a proteinaceous molecule or fragment thereof according to claim 11, an antigen according to claim 12, or an antibody according to claim 13 for the production of a pharmaceutical composition.
- 15 22. Use according to claim 22 for the production of a pharmaceutical composition for the treatment or prevention of an MPV infection.
  - 23. Use according to claim 22 or 23 for the production of a pharmaceutical composition for the treatment or prevention of respiratory tract illnesses.
  - 24. A pharmaceutical composition comprising a virus according to any one claims 1 to 7, a nucleic acid according to claim 8, a vector according to claim 9, a host cell according to claim 10, a proteinaceous molecule or fragment thereof according to claim 11, an antigen according to claim 12, or an antibody according to claim 13.
  - 25. A method for the treatment or prevention of an MPV infection comprising providing an individual with a pharmaceutical composition according to claim 24.
  - 26. A method for the treatment or prevention of a respiratory illness comprising providing an individual with a pharmaceutical composition according to claim 24.

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- 27. A method according to claim 25 or 26 wherein said individual comprises a human.
- 28. A method to obtain an antiviral agent useful in the treatment of
  respiratory tract illness comprising establishing a cell culture or experimental
  animal comprising a virus according to any one of claims 1 to 7, treating said
  culture or animal with an candidate antiviral agent, and determining the effect
  of said agent on said virus or its infection of said culture or animal.
- 10 29. An antiviral agent obtainable according to the method of claim 28.
  - 30. Use of an antiviral agent according to claim 29 for the preparation of a pharmaceutical composition.
- 15 31. Use according to claim 30 for the preparation of a pharmaceutical composition for the treatment of respiratory tract illness.
  - 32. Use according to claim 30 or 31 for the preparation of a pharmaceutical composition for the treatment of an MPV infection.
  - 33. A pharmaceutical composition comprising an antiviral agent according to claim 29.
- 34. A method for the treatment or prevention of an MPV infection comprising providing an individual with a pharmaceutical composition according to claim 33.
  - 35. A method for the treatment or prevention of a respiratory illness comprising providing an individual with a pharmaceutical composition according to claim 33.
  - 36. A method according to claim 34 or 35 wherein said individual comprises a human.

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**EPO-DG 1** 

Abstract

118, 10, 2001

The invention relates to the field of virology. The invention provides an isolated essentially mammalian negative-sense single stranded RNA virus (MPV) within the sub-familie *Pneumovirinae* of the family *Paramyxoviridae* and identifiable as phylogenetically corresponding to the genus *Metapneumovirus* and components thereof.

EPO - DG 1

(18. 10. 2001

TSIGAT	1



M							
	00-1	hRSV	brsv	PMV	APV-A	APV-C	APV-B
00-1	1,00	0,37	0,37	0,37	0,77	0,87	0,75
hRSV	~	1,00	0,91	0,41	0,37	0,37	0,37
brsv			1,00	0,42	0,35	0,36	0,35
PMV				1,00	0,37	0,38	0,38
APV-A					1,00	0,78	0,89
APV-C						1,00	0,77
APV-B							1,00
N							
•	00-1	hrsv	brsv	PVM	APV-A	APV-C	APV-B
00-1	1,00	0,20	0,22	0,21	0,40	0,52	0,40
hRSV		1,00	0,59	0,30	0,18	0,21	0,18
bRSV			1,00	0,31	0,21	0,23	0,21
PVM				1,00	0,21	0,23	0,21
APVA					1,00	0,42	1,00
APVC						1,00	0,42
APVB							1,00
F							
_	00-1	hRSV	brsv	PVM	APV-A	APV-C	APV-B
00-1	1,00	0,32	0,33	0,37	0,67	0,80	0,66
hRSV		1,00	0,82	0,40	0,35	0,35	0,35
bRSV			1,00	0,41	0,34	0,36	0,34
PVM				1,00	0,38	0,38	0,39
APV-A					1,00	0,72	0,84
APV-C						1,00	0,72
APV-B					·		1,00
P							
	00-1	hRSV	bRSV	PMV	APV-A	APV-C	
00-1	1,00	0,25	0,26	0,27	0,55	0,67	
hRSV		1,00	0,81	0,30	0,28	0,26	
bRSV			1,00	0,29	0,28	0,26	
PMV				1,00	0,23	0,27	
APV-A					1,00	0,52	
APV-C						1,00	
T8							
	00-1	hRSV	bRSV	APV-A			
00-1	1,00	0,36	0,35	0,56			
hRSV		1,00	0,79	0,36			
bRSV			1,00	0,35			
APV-A				1,00			
L9/10							
	00-1	hrsv	brsv	APV-A			
00-1	1,00	0,30	0,30	0,53			
hRSV		1,00	0,83	0,34			
bRSV			1,00	0,34			
APV-A				1,00		•	
··· v · A				1,00			

Fig. 1a

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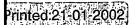
Table 2

Seroprevalence of hMPV in humans categorised by age group using immunofluorescence and virus neutralisation assays

	Immunoflu	orescence assays	Virus neut	ralisation assays	3
Age (Years)	N tested	N positive	N tested	N positive	Titre range
< 1	20	5	12	3	16-32
1-2	20	11	13	4	. 16-32
2-5	20	14	8	3	16-512
5-10	20	20	4	4	32-256
10-20	20	20	4	3	32-128
> 20	20	20	4	3	32-128
8-99 <sup>1</sup>	72	72	11	11	16-128

<sup>&</sup>lt;sup>1</sup>Sero-archeological analysis using sera collected in 1958

Fig. 1b





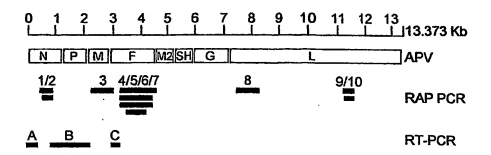


Fig. 2

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Fig. 3

#### Nucleo protein

```
00-1 NP MSLQGIHLSDLSYKHAILKESQYTIKRDVGTTTAVTPSSLQQEITLLCGEILYAKHADYKYAAEIGIQYISTALGSERVQQILRNSGSEVQVVLTRTYSL 10
     ...ES.R....E.....ED.....R...A...I...E..PQVST...MV.F...T..EP...V.M.......AD.T....K......G.M.KIVT. 10
     APV C
      .A.SKVK.N.TFN.DQL.ST.K...Q.ST.DNIDIPNYDV.KHLNK...ML.ITED.NH.FTGL..ML.AMSR..R.DTIK..KDA.YQ.RANGVDVITH 10
hrsv
      .A.SKVK.N.TIN.OQL.SS.K...Q.ST.DNIDTPNYDV.KHLNK...ML.ITED.NH.FTGL..ML.AMSR..R.DYIK..KDA.YH.KANGVDITTY 10
hRSV
      ...DRIK.N.V.N.DSL.SNCK.SVT.ST.DV.S.SGHAM.KALARTL.MF.LTAFNRCEEV....L..AMSL..RDDSIK...EA.YN.KC.D.QLKDF 10
PVM
00-1 NP GKIKNNKGEDLOMIDIHGVEKSWVERIDKEARKTMATLIKESSGNIPONORPSAPDTPIILLCVGALIFTKLASTIEVGLETTVRRANRVLSDALKRYPR 20
     SAEGSVRKREV--.N..D.GVG.ADDVERTT.EA.GAMVR.KV-QLTK.K..L.A.V...I.....V....AI...S.....IS....19
      APV C
      RQDV.G.EMKFEV.TLVSLTSEVQGN.EI.S..SYKM...M-.EVAPEY.HDS..CGM.V...A..VI....AGDRS..TAVI....N..RNEM...KG 19
brsv
      RQDI.G.EMKFEV.TLSSLTSEIQVN.EI.S..SYKM...M-.EVAPEY.HDS..CGM.I..IA..VI....AGDRS..TAVI....N..RNEI...KG 19
hrsv
      TIKLQG.EYKI.V...V.IDAANLADLEIQ..GVV.KE..TG-ARL.D.R.HD...CGV.V...IA..VVS...AGDRG..DAVE...IN..KAEKA...N 19
00-1 NP MDIPKIARSFYDLFEQKVYHRSLFIEYGKALGSSSTGSKAESLFVNIFNQAYGAGQTMIR#GVIARSSNNIMLGHVSVQAELKQVTEVYDLVRENGPESG 30
     ...R. K. FE. K. Y.N. T. RN. 29
APV A
APV B
      APV C
      LIPKD. N. .. EV. . KYPHYIDV. VHF.I.QS. TRG. . RV.GI. AGL. N. ...... V. ..... L.K. VK. ..... A. ... ME. . V. ... EYAQKL.G.A. 29
brsv
      LIPKD. N. .. EV. .KHPHLIDV.VHF.I.QS.TRG..RV.GI.AGL.N. ..S..V. .... L.K.VK. .... A. ... ME..V. .. EYAQKL.G.A. 29
hRSV
      .EVKQ..E.....R.P.YIDV..TF.L.QS.VRG...V.G..SGL.N....V...LL.K.VK....A...ME..V...EYAQKQ.G.A. 29
PVM
00-1 NP LLHLRQSPKAGLLSLANCPNFASVVLGNASGLGI IGMYRGRVPNTELFSAAESYAKSLKESNIKINFSSLGLTDEEKEAAEHFLNVSDDSQNDYE
     .....T....A......K..A..L...A.....RT.R.N...LAA....D.R...TSY.GGD.ERSSKF.
APV A
      .....N....E....N...INEEG....
APV B
      .....N. ......E. ....N. INEEG. ...
APV C
      FY.ILNN...S....TQF...S......A...M.E...TPR.QD.YD..KA..EQ...NGV..Y.V.D..T..L..KNQ..PK.N--DVEL
                                                                                 39
brsv
      FY.ILMN...S....TQF...S.....A...M.E...TPR.QD.YD..KA..EQ...NGV..Y.V,D..A..L..IKNQ..PKE.--DVEL
                                                                                 39
hRSV
      FY.I.NN...S...T.....T.....A.....S.K.APR.R...D.KD.ER.DN.V.Y.A.N.A.R.LISQQ.IV.TPD.DI
```

#### Phospho protein

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0-1 P	MS-FPEGKDILFMGNEAAKLAEAFQKSLRKPGHKRSQSIIGEKVNTVSETLELPTISRPAKPTIPSEPKLAFTDKGGATKTEIKQAIKVMDP	91
PV-A	M.D.YRNTSAG-GRSFIIA.KVPPLCN.TTSCI.PNKAPVPK	76
PV-C	LRRK.IR.TV.D.IIV.KKST.V.T. P.R.NGE_PDT_RSQTEE_RNEAT.	91
RSV		80
rsv	.EK.APEH.ED.NNK.TK.LES	45
VM	.EK.APEV.ED.N.KE.LKHRSFPSE.P.AGIPNTATHVTKYNMPPILRSSFKSPRVA.NL.EAPTTPPP.PPQN.EEQPKESD	92
0-1 P	IEEEESTEKKVLPSSDGKTPAEKKIKPSTNTKKNVSFTPNEPGKYTKLEKDALDLLSDNEEEDAE-SSILTFEERDTSSLSIEARLESIE	18
PV-A	IIYP.LPTAPVATDTYTSTSTE.AKKSK.DNPKVEEG.EPDND.KKA.TA	16
PV-C	EDASRLY.EVFA.TGKETPEKPT.KND.SRMEEDDVKALD	18
rsv		16
RSV	.DI.VTK.SPITSGTNIIN.TSEADSTPETKANYPR.PLKEDLTPSDNPFSY.ETIETFDNNEE.SYSYINDQNDN.TDR.D	13
VM	VDI.TMHVCPDNPEHSKKPCCSDDTD.KKTR.PM.T.VEP.EKFV.LGAS.YRETMQTFAADGYDEE.N.STNQEPG.S.V.QDR	18
0-1 P	EKISMIIGILRTINIATAGPTAARDGIRDAMIGVREELIADIIKEAKGKAAEMMEEEMXQRSKIGNGSVKITEKAKEINKIVEDESTSGESEGE	27
PV-A	M.KMMS.MTD.IKDTADLQ.SS.	25
PV-C	VV.L	27
RSV	x.	26
RSV	EM.HVV.SSV.LM.EK.RALATNORLEA.ARLRNSEKMA.DTSDE.P.NPTS.K.SDLLN	23
W	Y.IN.IMVTELTEM.KSDILTVNDRIVA.EKLRDCSRADTDDGSACYDR.RI.DSSNA	27
0-1 P	EEPKDTQDNSQEDDIYQLIM.	29
FV-A	SGESESDEE.SNLDLL	28
PV-C	EE.EEESNPDL.SLTMLIKN	29
RSV		28
RSV	SDNDLSLDF.	24
WW	EEAKEDLDVMGINFLI	29







### Matrix protein

## Fig. 3, contd.

```
00-1 malrix MESYLVDTYQGIPYTAAVQVDLIEKDLLPASLTIWFPLFQANTPPAVLLDQLKTLTTTLYAASQNGPILKWASAQGAAMFVLPKKFEVNATVAXDEYS 10
        ....II.....V......V....NN. .K..V......SS. .AP......S...Q.TV.PE..V.Q...T....SA.....S.S.AA.L.... 10
        ....II.....V...........SN..T..V......SS..AP.......S...Q.T..FE..V.Q...A.....SA.....A.S.A..L.... 10
APV-A
        APV-C
        ..T.VNKLHE.ST.....YNV....DD.....V.M..SSISADL.IKE,INVN.LVRQISTLK..S..IMINSRS.VLAQM.S..TIS.N.SL..R. 10
hRSV
        ..T. VNKLHE.ST.....YNVL...DD.....V.M. SSV.ADL. IKE ASIN. LVKQISTPK...S.R. TINSRS. VLAQM. SN. IIS.N. SL.. R. 10
hrsv
        ..A. ..EM.H.V. .......IN.V. ..HSANI ...V.I.M. .TSL .KNSVM.L. HDV.VICTQISTVH ..MI..DL.SSN.GLATM.RQ.LI..II.L.DRG 10
00-1 malrix KLEFDKLTVCEVKTVYLTTMKPYGMVSKFVSSAKSVGKKTHDLIALCDFMDLEKNTPVTIPAFIKSVSIKESESATVERAISSEADQALTQAKIAPYRGL 20
        ..D.GV...D.RA....L.....I.TMMNT..R.....I.M.RGI.....Y..A...D...........G....I..R...... 20
APV-A
        APV-C
        ..AY.IT.P..I.ACS..CL.VKN.LTTVKDLTMKTFNP..EI....E.ENIMTSKR.V..T.IR.INV.AKOLDSL.NIATT.FKN.I.N...I..... 20
brsv
        ..AY.VT.P..I.ACS..CL.VKS.LTTVKDLTMKTFNP..EI....E.ENIMTSKR.I..TYIRPI.V.NKDLNSL.NIATT.FKN.I.N...I..... 20
hRSV
        00-1 matrix IMINTMNPKGIFKKLGAGTQVIVELGAYVQAESISKICKTWSHQGTRYVLKSR.
APV-B
        25
APV-A
        27
APV-C
         HRSV
        VLVI.VTDN..A..YIKPQS.F..D...LEK...YYVTTN.K.TA.KFSI.P-
                                                                              25
hRSV
        VLVI.VTDN..A..YIKPQS.F..D...LEK...YYVTTN.K.TA..FSI.P-
        TLVINITST. .A. .L.K. .S. ILA. ..P.LTQV.LHDVIMN.K.T. .S. I ...SS---
```

#### Fusion protein

```
00-1 F MSW---KVVIXFSLLI-----TPXHGLKESYLFESCSTITEGYLSVLRTGWYTNVFTLEVGDVENLTCADGPS--LIKTELDLTKSALRELRTVSADQ 88
.ATTAMRMI.SIIFISTYVTHI.LCQNIT.EFYQST..AVSR....A......S.V.I.LSKIQKNV.KSTD.KVK...Q..ERYNN.VV..QSIMQNE 10
brsv
     .ELLIHRLSAI.LT.AINALYL.SSQNIT.EFYQST..AVSR..F.A.....S.I.I.LSNIKETK.NGTDTKVK...Q...KY.N.VT..QLIMQNT 10
        -- IPGR. FLV. . VIFNTKPIHPNT. T. K.Y. ST. . VE.A. . K.A. . . . . HMT . MSIKLSQINIES . KSSN. -- . LAH. . AIYS . . VD. . . . L. SNA 93
                        ----ienprosrfylgaialgvataaavtagvalaktirlesevtaiknalkktneavstlongvrvlatavrelk 16
00-1 F LAREEO-
            L......G..K.....RN......ND.. 16
APV-A V.K.SR---
APV-B ITK.NR-
                        ---.LSH.KK......T.....L......G..K...L..RS......I....ND.. 16
                          P.SFSRAKRGIPELIHYTRNSTKKFYGLMGKK.KR..L--GFL..IG--S..AS...VS.VLH..G..NK....LS..K..VS.S...S..TSK.LD.. 19
     P. ANNRARREAPOYMYTINTKNLINVS.SKK.KR..L--GFL...G--S.IAS.I.VS.VLH..G..NK.....IS..K..VS.S...S..TSK.LD.. 19
                          -LKSK.KK..L--GLI..LG--.....L...VQ....IAL.RD.VRN.....VS.T..MS...KV.DD.. 16
00-1 f devsknlirainknkodiadikmavsesgenrefinvvrgesdnagitpaisloimtdaelaravsnmptsaggikimienramvrrkgegeligvygss 26
APV-A E.I.K.P.Q.N.I.I.G.N.S.S.V.D.V.INR.S.S.N.D.V.D.D.V.ORT 26
APV-B E.I.K.P.Q.N.IR.J.G.N.S.S.V.VK.INR.S.S.N.
     NYID.E. LPQV. NHD.R. SNIETVIE.Q.K.N.L.EIA.E..V.....TPL. TYML.NS..LSLIND..ITND.K...SS.VQI..QQSYSIMSV.KEEV 29
     NYINNQ.LPIV.QQS.R.SNIETVIE.Q.K.S.L.EIN.E..V...V.TPL.TYML.NS..LSLIND..ITND.K...SS.VQI..QQSYSIMSIIKEEV 29
     N.I..E.LPK..RVS..VH.ITAVIR.Q.L.K.L.E.S.E..S...L.HTV.SFML.R.TSI.GG.AV...KEI..SSK.IM..N.LAI.SS.NADT 26
00-1 F VIXMVQLPIFGVIDTPCWIVKAAPSC---SGKKGNYACLLREDQGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSXECNINISTTNYPCKVS 36
APV-A .VY.....E....R.V...L.--RKE.....I......T....A....KD...V.D.Y....LEVEQ..Y...SK..... 36
    .VY.....E....R.V...L.--RHERES.....T....A....D...V.D.Y......SEVEQ..H....ST...... 36
     IAYV....Y......KLHTS.L.TTDN.E.SNI..T.T.R...D....VSFF.QTET.KVQSNR.....MNSLTLPTDVNL..TD.FN.K.D..IM 39
     LAYV.....Y......KLHTS.L.TTNI.E.SNI..T.T.R...D....VSFF.QADT.KVQSNR.....MNSLTLPSEVSL..TD.FNSK.D..IM 39
     LVYVI...L..M.D..VIRSSID.--HNIADK....A.A.N...H...LS.F.SPT...IHNGYA...LKSLT.PVF.R...S.MY....D..I. 35
00-1 F TGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCSYITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDFVKFPEDQFNVALDQVF 46
.SKTD..SSVITSI..I.S..GKTK.TASNKORR....TFSN..D.VS.KGV...SVG..L.YVN.L..KALY...E.IIMYY..LV..S.E.DASIA..N 49
     .SKTD..SSVITS...I.S..GKTK.TASNKNR....TFSN..D.VS.KGV...SVG..L.YVN.L..KNLYV..E.ILNYY..LV..S.E.DASIS..N 49
     .SKTYV.TAV.TTM.C..S..GHN., TVIN.DK...RT.PD..H..S.KGV.R.QVG....Y...EV.KSI.VR.E.LVLKY..LS..D.K.D..IRD.E 45
00-1 F ESIENSQALVDQSNRILS----SAEKGNTGFII--VIILIAVLGSTMILVSVFIIIKKTKRPTGAP-PELSGVTNNGFIPH-N.
APV-A ...DR..D.I.K..DL.G----ADA.SKA.IA.--A.VVLVI..IFFL.AVIYYCSRVR.TKPKHDY.ATT.HSSMAYV--
APV-B ..VDK.KD.I.K..DL.D----IEV.S.I.AAL--A.TILV..SMLI.VGIAYYVV..R.AK.SNGY.KTT.QS.M.Y.S.
                                                                                53
    ..V.K..N.I....K..D----.I....A..V.--..V..VI.MIAAVG.G..FVV..R.AAPKF.-M.MN..N.K...--F.LIKKKKKKKKK
    AK.NQ.L.FIRR.DEL.H--SVDVG.ST.NVV.TTI..V.V.VIIML.A.GLLFYC.TKST.IMLGKDQ...IN.LS.S--
brsv
                                                                                57
     .K.NQ.L. FIRR.DEL.H--NVNTG.ST.NIM.TTI..V.I.VILSL.AIGILLYC.AKNT.VTISKDQ...IN.IA.S--
hRSV
                                                                                57
     H...NQTRTFFKA.DQL.DLSENREN.NLNKSY.LTTLLFVVM.III.AVIGFILYKVLK----MIRDNK.KSKSTP.LT-
```

hrsv

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# Fig. 3, contd.

## L polymerase RAP PCR fragment 8

APV-A ME-IS brsv MDTL hrsv MDPI	TYNVYLPDSYLKGVISFSETRAIGSCLLKRPYLKNONTAKVALENFVIENVRAVNSKERLSJYK	90 90
APV-A LL brsv L.TY	KNVHSCELTLLKQFLTRSKNISTIKINMICDWLQIKSTSDDTSILSFIDVEFI	17
APV-A brsv pdop		20
AFV-A brsv	WMLSRENANECIWVSNSLNENQEGVGLRSNLQ LAL.VCSA.D.LK.VGELLAR ISL.VCMIT.ICTLAKSLC ISL.VCLIT,ICTLAKSL.	2: 3: 3:
L polymerase R	AP-PCR fragment 9/10	
00-1 fragment 9/10 APV-A bRSV hRSV	KLVDKITSDCHIFSPDKIDMLTLGKMLMPTIKGCKTDQFINKRENYFHGNNLIESLSAALAXHWCGILTEQC -F.S.RVTN.HLVML.LVRSNINNN	7:
00-1 fragment 9/10 APV-A bRSV hRSV	IENNIFKKDWGDGFISDHAFMDFRIFLCVFKTKLLC  T.S.Q.ETIN.TWMSYHW  DSKG.EE.Y.TM.L.LAV.FDAYY.  DSKG.EE.Y.TM.INL.V.FNAYY	1 1 1

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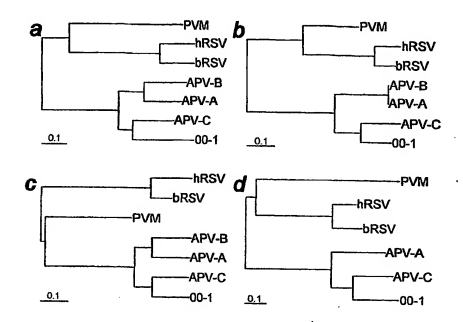


Fig. 4

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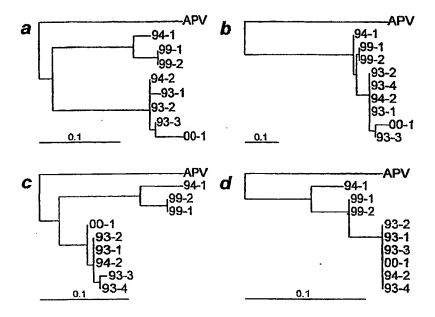


Fig. 5

ICGCGIA IAAATIAGAI ICAAAAATATGGGACAAGTGAAAATGIC ICTICAAGGGATTCACCTGAGIGATTAICATACAAGCAIGEIATAIAAAAGAGTCTCAGTACAAAAAAAAAAAGAGATGIGGGTACAACAACTGCAGTGACA

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Fig. 6A

9 288 1350 8 ဋ္ဌ 8 750 8 505 58 쯇 1950 accitgcttaabgaatcatcagaaaatcagabggccctcagcaccagalgccctcataatctattatbtbtabgtgccttaatattcactaakctagcatcaagaagtgggactagagccacagtcagagg gctaaccgtgtactaagtgatgcactcaagagataccctagaatggataccaaagattgccagatccttgtatgactaatagagagaatatggtaatgagtaaggcaaagcattagcaatcatcatca GBCABCAAABCAGAAABICTAIITGITAATATTCATGCAAGCITATGGGGCCGGTCAAAGATGCIAAGGTGGGGGGTCAITGCAGGTCATCCAACAATATATGITAGGACAIGTATCGGTCCAAGC1GAGTTAAAACAGGTCACA gartelatbactiggtggagaaatgggccctgaatetggacttetacattaaggccaaagccgaaagctgattgttagcgaactgtcccaaetttgcaabtgtctcggaaatgcctcaggcttaggcataatgg y giat carggaaga taratara aataattatti tagcaactigaaactiat gccaaaaqttigaaagaaataaataaatii cittattagaattagaattagag GACGACAGICAAAATGATTATGAGTAAAAAAGIGGGACAAGICGAAAATGICCTGAAGGAAAAGATTCTTTICAIGGGTAAGAAGGAGCAGAAAATAGCAGAAGCTTICCAGAAATCATTAAGAAAACCAGGICATAAAA aagcaatcaabtcatggatccaitgaagaagaagttaccgagaagaggtsctacctccagtgatgggaaaccctgcagaaagaaaggaaccaigaaccaaaaagaaggtitcatttackcaaatgaaccag sbaatatacaabttobaaaabatgctctagatitgctctcagataatbaagaagatgcagaatcttcatcttaaccittgabaaagaatacttcatcattaagcattgaggccagattggaatcatagggaaaattaa EIOKEARKT N K I N F S S L G L T D E E K E A A E H F L N ы MOAY GAGOTHLRK GYIARSSKNIHLGHV SSLOOEITLLCGEILYAKHADYKYAAEIGIOYISTA တ N R V L S D A L K R Y P R M D I P K 1 A R S F Y D L F E Q K V Y H R S LYREMGPESGILHIROSPKAGLISLANCPNFA DAESSILTFE.ERDISS G G A L I F NIATAGPTAAROGIRDAMIGVRE H - 0 - H 0 E KSFPEGKOIL EKKYLPSSDG N K G E D KIGNGSVKLTEKAK SOSIIGEKVNIVSETLELPII 8 7 z EEST YPNTELFSA DAIKVNDPI တ S

GTCAAGAAGAIGACAIIIACCAGIIAATIAIGIAGIIIAATAAAAAIAAAGAATGGGACAAGTAAAAATGGABICCIACCTAGGAGACCTATCAAGGCAITCCITACACAGCAGCTGTICAAGTIGATGTAAAAAAGGACCTGTI

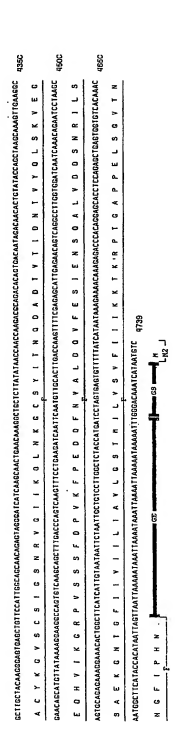
10/23

Fig. 6A, contd.

	YFCDIAAGINVAEDS?ECHINISTINYPCKYSIGRHPISHYALSPLGALV
420C	GTCTTTTGCGACACAGCAGGAATCAATGTTGCTGAGGAGTGCAARGGAGGGGACATAAACATATCTACTACTACCATGCAAAGTTAGCACAGGAAGACATCCTATCAGTATGGTTGCTTGC
	CHIVKAAPSCSGKKGNYACLLREDOGWYCONAGSTVYPNEKOCETRGOM
4050	IGCTGGATAGTAAAABGAGCCCCTTCTTGTTCAGGAAAAAAGGGAAACTATGCTTGCCTCTTAAGAGAAGACCAAGGATGGTATTGTCAAAAGGGTGAACTGTTACTACCAAATGAAAAAAAA
	NHPTSA601KLNLENRANVRRKGFGFLIGVYGSSVI?NVOLPIFGVIOTP
3900	AACAIGCCAACAICIGCAGGACAAAIAAAACTGAIGIIGGAGAACCGIGCAAIGGIAAGAAAGAAAAAAAAA
	O I A O L K B A Y S F S O F N R R F L N Y Y R O F S O N A G I T P A I S L O L M T O A E L A R A Y S
3750	GACATIGCTGACCTGAAAATGGCCGTTAGCTTCAGTCAATTCAACAGAAGGTTCCTAAATGTTGTGCGGCAATTTCGGAATAACACCAGCAATATCTTTGGACTTAATGACAGAAGCTGAACTAGCCAGAAGCTGTTCC
	E Y T A I K H A L K T N E A Y S I L G N G Y R Y L A T A Y R E L K D F Y S K N L T R A I N K N K C
3800	GAAGTAACAGCAA1TAAGAATGCCCTCAAAAAAGACCAATGAAGCAGTATCTACATTGGGGAATGGAGTTCGTGTGTGGCAACTGCAGGAGCTGAAAAAAAA
	TVS A D O LAREEO IENPROSRFYLGA I ALGVATAAAVTAGVAJAKTIRLES
3450	ACAGITICIGATCAALTBBCAABABGAGGAAATTBAAAATCCCAGACAATCIAGATTCGTGTGTGTGTGTAGCACTCGGTGTTGCAACTGCAGCTGTTACAGGTGTTGCAATTGCAAAACCATCGGCTTGAAAGT
	E GYLSYLRTGWYTNVFTLEVGDVENLTCADGPSLIKTELDLTKSALRELR
3300	GAAGGATATCICAGTGTTCTGAGGACAGGTTGGTTGGTTTTTACACTGGAGGTAGGCGATGTAGAGAACCTTACATGTGCGACCTAGACTAGACTGAGAATTAGACCTGACCAAAAGTGCACTAAGAAGAATTAGACCTGACAAAAGTGCACTAAAGAAGAATTAGACCTGACAAAAGTGCACTAAAGAAGAATTAGACTGACAAAAGTGCACTAAAGAAGAATTAGACTGACAAAAGAATTAGACTGAAAAGAAATAAAAAAAA
	GP H S W K V V I 7 F S L L I T P 2 H G L K E S Y L E E S C S I 1 I
3150	ATAAAAATCAACTTAGAACAAGAATTAAATCAAGAACGGGACAAATAAAATAAAATGCTTGGAAGTGGTGATCAKTTTTCATTGTTAATAACACCTCRACACGGTCTTAAAGAGGGCTACTTAGAAGAGCTACTTAGAAGAGCTACTTAGAAGAGCTACTTAGAAGAGCTACTTAGAAGAGCTACTAGAAGTGTATAAAGT
	Y O A E S I S K I C K T W S H O G T R Y Y L K S R .
3000	ATAIGICEAGGCIGAAAGCATAAGCAAAATAIGCAAGACTIGGAGCCAICAAGGGGACAAGAAGATTGIGICIIGAAGICCAAACAACCAAGCACCIIGGGCAAGAGCTACTAAGCTAICICAIAGAICAIAAGGCACCAAAGAGAGAAAAAAAAAA
	AAISSEADOALTOÄKIAPYAGLIHIHIHNNFKGIFKKLGAGTOVIVELGA
2850	AGCIGCTATAAGCAGIGAAGCAAGCTGTAACAGGCCGAAAATIGCACCTTAIGCGGGATTAATTATGATCAIGACTAIGAACAATCCCAAAGGCATATTCAAAAAGCTTGBAGCTGGGGCTCAAGTGATAGTAGTAGGAGC
	FYSSAKSYGKKTHDLIALCOFMDLEKNTPYIIPAFIKSVSIKESESATVE
2002	ATTIGIGAGCICAGCCAAATCAGTIGGCAAAAAAAACACATGATGTAATCGCACTATGTGATTTTATGGATGTAGAAAGAA
	AAHFYLPKKFEYNATVA7DEYSKLEFDKLTYCEYKTYYLTIMKPYGHYSK
255C	TECAGEAATGITTGTACTICCCAAAAAAITIBAAGTGAATGCGACTGTAGCANTCGATGAAATATAGCAAACTGGAAATTTGACAAACCAGTGTGAAAAAAAA
	PASLTIMFPLFOANTPPAVLLOOLKTLTITTYAASONGPILKVNASAOG
240c	ACCIGCAACCCTAACAATAGGITCCCTTTGTTTCAGGCCAACACCACCACCAGCAGCGCTCGATCAGCTAAAAACCCTGACAATAACCACTGTATGCTGAAAATGGTCCAAAATGGTCAAAAGTGAATGCTCAAAGTGAATGCTCAAAGTGAATGCTCAAGGTGAATGCTCAAGGTGAATGCTCAAGGTGAGCTCAAGG
	SOEODIYOLIM.

11/23

Fig. 6A, contd.



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CAAGAAAAAALIGIICCACIGIIAAIGICIACIICIIGACICATAICIIGAGAGGAGIGATIICCIIIAGIGAGACTAAIGCATIGGIICAIGIICTITAAAAAACCTIACCAAAAAAITGACAACAGIIGCAAAAAGIIGCAAAAA

aatctiitatcgagcattiagactcaaaaatgcagtcaattctaagatgaaatatcagattacaagatagtaggccagtaaacatgcaacataabaatgagcagttgtgag

O E K H C S T YNYY L P D SYLK G Y I S F S E T N A I G S C L L K R P Y L K N D N T A K Y A I

acaggagi aaaatititagcactotcaatitaaatatgatatgotatogatigggotgggottaaagtctacatgagatgatactctaabottaabititataggaatitatagggagtgsggaagtggottabtaatggsta

NP VIEHVRLKNAVNSKAK 1 SOYKIVEPVNADHE I AKNVHS CELTLLKOF

AAICICAACAAGIIGAIICIGGAAITCAGGAAAGAAGAAGIAAIAAGAACIGGIICAAICIIGIGIAGGICAIIGGSTAAIIAGIIIIIIGIIGIAIGAIAIGGAIGIAIAGICAAGAGAAAAAAGAGAAAAAGAIGAACIICIIC

1 R S K N I S 1 L K L N H I C D N L O L K S 1 S D D 1 S I L S F I D V E F 1 P S K V S N M F S N W Y

T T N O L L T W K D V H L S R F N A N F C J W V S M S L N E N O E G V G L R S N L O G I L T N K L Y

781

GAAACTGTAGATTATATGCTTAGTTTATGTT

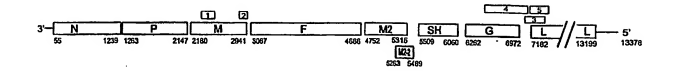
ETVOYBLSLC

N L N K L I L E F R . K E E Y 1 R 1 G S . L C R S L G K L Y F Y Y S S Y G C 1 Y K S N K S K R Y S F F

-ig. 6B

င္ည ద్ద AIAAGCTAGIAGATAAGATAZGATCTICTGATCATATCTTCAGTCCAGACAAAATAGATAIGTTAACACTIGGGGAAAAATGCTCATGCCCACTATAAAAGGTCAGAAAAACAGATCAGTTCTGAACAAGAGAAATAAGTTCCTAAGGA N N L I E S L S A A L A C H W C G I L T E O C I E N N I F K K O W G D G F I S D H A F M D E K I F L DOHIFSPDKI DRITLEKALAPTIKOOKTDOFLNKRENY GIGICITIAAAACIAAACITITAIGIA 327 K L V O K 1 T S C Y F X T K L L C Fig. 6C





```
MSLQGIHLSDLSYKHAILKESQYTIKROVGTTTAVTPSSLQQEITLLCGE
hMPV
       APVC
APVA
hreva
brsv
        ...DRLK.N.V.N.DSL.SNCK.SVT.ST.DV.S.SGHAM.KALARTL.M
PVM
        ILYAKHADYKYAAEIGIQYISTALGSERVQQILRNSGSEVQVVLTRTYSL
MOV
        ....T.SH..V.M..V.T.A.T..K..A..K...
F. EP. QV.M...ADKT..KS...G.M.KIVT.
V.F..T.EP..V.M...AD.T..K...G.M.KIVT.
APVC
APVB
APVA
        L. ITED. NH. FTGL. ML. AMSR. R. DTIK. DA. YH. KANGV DYTH
L. ITED. NH. FTGL. ML. AMSR. R. DTIK. KDA. YH. KANGV DITTY
hreva
        L. ITED.NH. FYGL. ML. AMSR. R. DTLK. KDA. YO. RANGY DYITH F. LTAFNRCEEV. L. AMSL. RDDSIK. . EA. YN. KC. D. QLKDF
hrsv
PVX
        GKIKNNKGEDLOMLDIHGVEKSWVEEIDKEARKTMATLLKESSGNIPONO
hMPV
        .G.S.E....R.I.V....SAT.DN.P...
PAEGFIR-KREV.N.DIGPA.ADNVERT.E.SLAV.K-AQ.K.
APVC
APVB
        SAEGSVR--KREV.N..D.GVG.ADDVERTT.EA.GAMVR.K-VQLTK.
APVA
        RQDI.G.EMKFEV.TLASLTTEIQIN.BI.S..SYKKM...M-.EVAPEY
RQDI.G.EMKFEV.TLSSLTSEIQVN.BI.S..SYKKM...M-.EVAPEY
RQDV.G.EMKFEV.TLVSLTSEVQGN.BI.S..SYKKM...M-.EVAPEY
hreva
LREVB
brev
        TIKLOG.EYKI.V...V.IDAANLADLEIQ..GVV.KE..TG-ARL.D.R
MVC
        A 20
RPSAPDTPILLCYGALIFTKIASTIEVGLETTVRRANKVLSDALKRYPR
HMDV
                    ...i....v....A.....N.....P...
         ...S..A.
APVC
 APVB
        K...L.A.V
        HDS. CGM I IA. VI . AGDRS. TAVI . N. KNEM . KG
HDS. CGM I IA. VI . AGDRS. TAVI . N. KNEI . KG
HDS. CGM V . A. VI . AGDRS. TAVI N. KNEM . KG
HD . CGM V . IA. VVS . AGDRG . DAVE . IN . KAEKA . N
 APVA
LRSVA
 hrsve
 HERV
 PVM
        MDIPKIARSFYDLFEORVYHRSLFIEYGRALGSSSTGSKA
 HMPV
        APVC
 APVA
 HREVA
 LRSVB
         LIPKD..N...EV..KYPHYIDV.VHF.I.QS.TRG..RV.GI.AGL..N
 brav
         PVX
         ), ere egi der litter fregenskripp at hør litterette flytter fregenskrip) af til etter sett.
 hMPV
 APVC
                                             ....R..8......K.....
 APVB
               R. S. K.
V. L. K. VK. A. ME. V. EYAQKI. G. A.
V. L. K. VK. A. ME. V. EYAQKI. G. A.
V. L. K. VK. A. ME. V. EYAQKI. G. A.
V. L. K. VK. A. ME. V. EYAQKI. G. A.
V. LL K. VK. A. ME. V. EYAQKI. G. A.
 APVA
 hRSVX
 hrsve
 brev
 DVM
         gregorium, Comesquenticio entre esponere de les RVPN CEMPSA
 MIPV
 FX.I.NN..S...T....T....A....S.K.APR.R...D.
        AESYAKSLKESNKINFSSLGLTDEEKEAAEHFLNVSDDS-QNDYE
 hMPV
         R.....E...N..INEEG-....R....R...LAA...ED.R...TSY.GGDS.K-SQKF.
  APVA
  APVA ....RT.R.N...LAA....D.R...TSY.GGD.RR-SSKF.

hrsva .KA.EQ...NGV.Y.V.D.A.L..IK.Q..PK.N-DVEL-

hrsvb .KA.EQ...NGV.Y.V.D.A.L..IK.Q..PKE.--DVEL-
  brsv
         .KA..EQ...NGV..Y.V.D..T..L..IKNQ..PK.H--DVEL-
          .KD. .ER. .DN.V.,Y.A.N..A..R.LISQQ..IV..TPDD.I-
```

```
MSFPEGKDILFMGNEAAKLAEAFQKSLRKPGHKRS-----QSIIGEK
 HMPV
        APVA
 APVB
 APVA
           --M.KFAPE.H.ED.NNR.TK.LE.----
brsvb ---m.kfape.H.ed.nnk.tk.le.-----brsvb ---m.kfape.H.ed.ntk.tk.le.-----
        ---M.KFAPE.V.ED.N.K..E.L.HRSF.SE.PLAGIPNTATHVTKYNM
 PVM
        VNTVSETLELPTISRPAKPTIPSEPKLAHTDKGGATKTEIKQAIKVMDPI
 HMPV
        APVC
 APVB
 APVA
        I..IA.KVP..PLCN.TT.-----..SCI.PN-.APVPKVK--
           -- IKGKFTS.-
                                    -----KDPKK.DS.ISVNS.
hrsva
        --- IKGKFASS-
 hrsve
                           ------KDPXK.DS.ISVNS.
         ---LKGKFTSS-
brsv
                                    ------KDSRK.DS.ISVNSV
        PPILRSSFK..SPRVA.NL.E..A.P---TTPPP.PPQN.EEQPKESDV
 PVM
MPV
        eeeestekkvlpssdgktpaekklkpstntkkk----vsftpnepgkyt
DASRLY.EVFA.T......GRETPEKP...---.K.ND.S.R.
APVE V.SIYP.LPTAPP.ANIETAHPIGAPKKAQ.R----.K. BSSKA...
AFVA I.SIYP.LPTAPVATD.YTSTSTESAKKS..----K. DNFKV...
BRSVA DI.VTK.SPITST.TIIN.THETODTAG.KPHYQRKPL..KEDLTPSUN
BRSVA DI.VTK.SPITSGTNIN.TSEADSTPETKANYPRKPL.KEDLTPSUN
DI.VTK.SPITSGTNIN.TSEADSTPETKANYPRKPL.KEDLTPSUN
DI.VTK.SPITSGTNIN.TSEADSTPETKANYPRKPL.KEDLTPSUN
hrsv
        DI.LPK.SPITSTNQNINQPSEINDTIATNQVHIRKPL...KEEL.SSEN
PVM
        DI.TMHVC..PDNPEHSKKPCCSDDTD.KKT---RKPM.T.VEP.EKFVG
hMPV
        KLEKDALDLLSD-NEEEDAESSILTFEERD--TSSLSIEARLESIEEKLS
APVC
        ..ME.E..-.DD...V...K.---A.L...D...
..EE.E...PD.ON.EK.V....K.--NAES....A...
APVA ...EE..E. PD.ON.EK..V...K.-NAES....A....APVA ...EEG.E. PE.DN.EK...K.--.A.T...A....A...ARVA PFS.LYKETIETFDNN--E.E.SYSY..INDQ.NIN-T...DR.D...ARSVA PFS.LYKETIETFDNN--E.E.SYSY..INDQ.NIN-T...DR.D...BRSV PFTRLYKETIETFDNN--E.E.SYSY..INDQ.NIN-T...DR.D...PVM LGASLYRETMQTFAADGYD.E.N.S...TNQEPG.S.V.Q.DR.....250
MOPV
        MILGLERTLNIATAGPTAARDGIRDAMIGVREELIADIIKEAKGK----
PVM
       Y.I. N.IMV.....T. E. L. T. EM KSDILTVNDRIV
                                                                      300
        -AAEMMEEEMSQRSKIGNGSVKLTEKAKELNRIV
hMPU
       APVC
       APVA
hrsva
breve AM.RLRN..SEKMA.DTSDE.P.NPTS.K.SDLL..---
brsv AM.RLRD..SEKMT.DTSDE...PTSEK..MVL..--
PVM AMEKLRD..C.RADTDDGSACY..DR.RI.D..SSNA-
                          316
            WINDTIAN DE LE CONTRE
hMPV
       .L.S.T.
...SFD.
S.A.S..N.DL
APVC
APVR
brsva
        .SDNDLSLE.F----
       .SDNDLSL..F--
brsva
       SSDNDLSLE.F---
       EAKEDLDV...MGINF
```

van den Hoogen et al., Figure

9

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50

MESYLVDT (QGIPTERNY VDLIEKDLLPASLTIWFPLFQANTPPAVLLD

. V. W. W. V. Q. R. V. V. T. T. E.

. II. V. S. N. K. V. SS. AP.

. II. V. SS. AP.

. T. VNKLHE. ST. YNVL. DD. V. M. SSV. ADL. IK

T. VNKLHE. ST. YNVL. DD. V. M. SSV. ADL. IK

T. VNKLHE. ST. YNVL. DD. V. M. SST. SADL. IK

T. VNKLHE. ST. YNVL. DD. V. M. SST. SADL. IK

A. EM. R. V. JELEVAL V. HSANI. V. I. M. TSL. KNSVM.
APV C
APV B
APV A
hrsv A
hrsv B
bRSV
PVM
                                                                                                                                                                                                   100
                           QLKTI.T ITT LYAASQNGPI LKVNASAQGAAMSVLPKKFEVNATVALDEYS
                         hmpv
APV C
APV B
 hRSV A
hrsv B
brsv
                            L.HDV.VICTQISTVH..MI..DL.SSN.GLATM.RQ.LI..II...DWG
 PVH
                            KLEFDKLTVCEVKTVYLTTMKPYGMVSKFVSSAKSVGKKTHDLIALCDFM
                           KLEFOKLTVCEVKTVYLTTMKPYGMVSKYVSSAKSVGKRIRDLIALUFM

LA. N. A. L.
D.GV. D.RA. L. I.TNMNT. R. I.
R. GT. D.RSI. L. INTDVR. R. I.
AY.VT.P. I.ACS. CL.SKN LTTVKDLTMKTLNP. I. E.E.
AY.VT.P. I.ACS. CL.VKS.LTTVKDLTMKTFNP. EI. E.E.
AY.IT.P. I.ACS. CL.VKN.LTTVKDLTMKTFNP. EV. VS. II.

AV. IT.P. I.ACS. CL.VKN.LTTVKNDTMKTTNP. EV. VS. II.

AV. IT.P. I.ACS. CL.VKN.LTTVKNDTMKTNP. EV. VS. II.

AV. IT.P. I.ACS. CL.VKN.LTTVKNDTMKTNP. EV. VS. II.

AV. IT.P. I.ACS. CL.VKN.LTTVKNDTMKTNP. EV. VS. II.

AV. IT. I.ACS. CL.VKN.LTTVKNDTMKTNP. EV. VS. II.

AV. I
APV C
  hrsv A
  hrsv B
  brsv
                             NMDYEVPVAFDK.SFCV.IL..KN.LYTVP.ITP-TNRP..E...V.S.H
  MVd
                                                                                                                                                                                                     200
                             DLEKNTPVTIPAFIKSVSIKESESATVEAAISSEADQALTQAKIAPYAGL
  hMPV
                            APV C
  APV B
  APV A
  hrsv A
  hrsv B
  hRSV
                             NRVTLKSFN..V..RALY.ROOGLDS..Q....DV.H.I.T.RV.....
  PVM
                              IMIMTMNNPKGIFKKLGAGTQVIVELGAYVQAESISKICKTWSKQGTRYV
                            hMPV
   APV C.
    APV B
    APV A
    hRSV B
    bRSV
    PVM
                               LKSR---- 258
    MPV
                               ...----
    APV C
                               ...---
    APV B
    APV A
                               I.PMED--
    hrsv A
                               I.PIED--
    bRSV
                               ...SSTSG
    PVM
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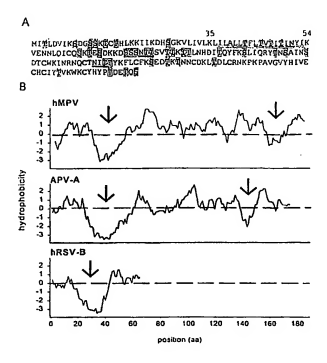


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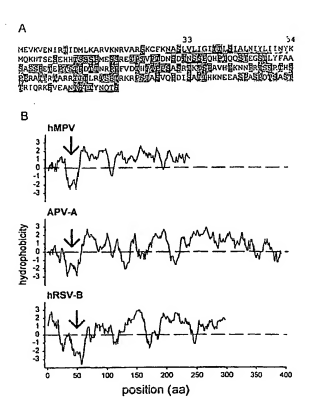
## 17/23

	919nal peptide  SECURITY TEST STATEMENT STATEM
hMPV	
APVC	
APVB	
APVA	
hRSVA	
hrsva	
brsv PVM	
PVM	MIPGRIFLVLLV. NTKPIHPNT.T.K.Y.ST. VE.A.K.AHMT.MSIKLSQINIES_KSSN,LAH.AIYSVDL.SN-
	And the Arms of th
hMPV	fusion domain 200
APVC	LAREEQ
APVB	. K.ARMS. KA
APVA	ITK.NR
PDCAV	V.K.SRLSS.RR. L. G.K.RN. ND. PPINNRARRELPRIMITINNAKKINVILSKK.KRFLGS.IASVS.VLH.G.NK.S.LS.K.VS.S.S.TSK.LD.
hRSVB	P.ANNRARREAPQYMYTINTKKLINVS.SKK.KRFLG-S.IAS.I.VS.VLHGNKLSKVS.SSTSK.LD
bRSV	P.SFSRAKRGIPELIHYTRNSTKKFYGLNGKK.KRFL.IGS.ASVS.VLH.G.NKLS.K.VS.SSTSK.LD.
PVM	
hMPV	DEVSK <u>NLI</u> RAINKNK DIAOLKMAVSESQENRRELNVVRQESDNAGITPAISLDIMTDAELARAVSNMPTSAGQIKLMLENRAMVRRKGEGELIGVYGSS
APVC	4 - 1 - 4 - 4 - 4 - 4 - 4 - 4 - 4 - 4 -
APVB	E-1NFUdillaIRIG-N
APVA	Palaghan Pangual Inggalaga Inggalaga Inggalaga Inggalaga Inggalaga Inggalaga Inggalaga Inggalaga Inggalaga Ing
hRSVA	NIIU.U.LPIVUS.B. SNIETVIE.U.K.N.IEIT.E. V. V. TRU TYMI NG. 1 CT YND. TOND W. AN AND
hrsve	
DV2A	NIAU-E-EFUV-NRIMIR-SNIAIVIE-U-N-N-L-EIR-EV TPI, TYMI, NS 1 CT TNN TWN P
PVM	N.I.E.LPKRVS.J.VH.ITAVIR.Q.L.K.L.E.S.ESL.HTV.9FML.RTSI.GG.AVKEISSK.IM.N.LAI.SS.NADT
hmpv Apvc	VIYMVOLPIFGVIDTFGWIVKAAPSGSGKKGNYAGLLREDOGWYGQNAGSTVYYPNEKIGETRGDHVIGDTAAGINVAEQSKEGNI <u>NIS</u> TTNYPFKVS
APVE	· · · · · · · · · · · · · · · · · · ·
APVA	.V E R.V LRHERES T A D V.D.Y
hRSVA	V. S. R.V. LRK-E. I. T. A. KD. V.D.Y SSVEG. H ST LAV. LY KD. V.D.Y LEVEG. Y SK LAV. LY KD. V.D.Y MNSLTLPSEINL. VD. FNPK. D IM
hRSVB	LA.VYKLHTS.L.TTNI.E.SNI.T.T.RDVSFF.QADT KVQSNRMNSLTLPSEVNL.VD.FNPK.DIM
brsv	
PVM	LV.VILMd.VIRSSIDHNIADKJ.A.A.NHLS.F.SPT.J.IHNGYA.JLKSLT.PVT.R.J.S.MYD.I.
<b>MPV</b>	TGRHPISMVALSPLGALVALIYKGVSDSIGSNRVGIIKOLNKG/ISVITNODADM/PTDNTUVOT CV//CCCO/INTY//CCC DV///CCCO/INTY//CCC
APVC	······································
APVB	
APVA	anna Vanna Tana Gan State Sauttana a Kanana a Gan DTH DINE T trong a company of
hRSVA	
hrsvb brsv	. SATU SSVITS I. S.J. GKTN. ITASNKNR TFSN. I.D. VS. KGV. SVG I. VIAI I. 101 VII. B. TTINGY T
PVM	SKTD. SSVITSII.S. GKTR TASNKNRTFSN. D.VS.KGVSVG. L.YVN.L.KALYE. IINYYLVS.E. DASISN
E 412	SKTYV.TAV.TIM.CS. GHN. TVIN.DKRT.PD. HS.KGV.R.QVGYEV.KSI.VR.E.LVLKY.LS. D.K.D. IRD.E
hMPV	ESIENSQALVDQSNRILSSAEKGMTG <u>NIIVDTETAVEGSTALTIVSVERIV</u> KKTKRGAPPELSGVTNNGFIPHN
APVC	.V.K.N.IK.D.IA.VV.LMLAAVG.GFVV.R.AAPKF.M.MN.N.K
APVB	VDK.KD.I.KDL.DIEVS.I.AALA.TILV.SMLI.VGIAYYVV.R.AKTSNGY.KTT.QS.H.Y.S
APVA	DRD.I.KDL.GADA,SKA.IA.A.VVLVI. IFPL AVIVYCERUR TKPK
hRSVA	.K.NO.L.FIRK.DEL.HNVNAG.ST.NIM.TT.I.VIIVILLS.IA_GLLLYCKA_STP-UTLSYDO IN TA SW
HRSVB	.K.NO.L.FIRR.DEL, HNVNTG.ST.NIM.TT.I.VIIVVLLS TATCLITYCKAKNTD-VTTICVDO TV TA CV
brsv	AK.NO.L.FIRR.DEL.H.VDVG.ST.NVV.TT.I.VIVVVIIM.IA.GLLFYCKTKSTP-TMLCKDO TM 10 CV-
PVM	H. NOTRIFFKA. DQL. DLS. NREN. NLNKSY. LTT. LF. VMLII. MAVIGF. LY. VIKMIRDNKKKSKSTP. L. VI.S

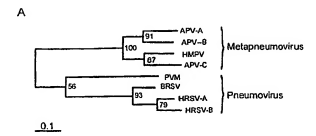
```
Α
          Kerkaperveyrekereseertuhnynsweurtealtertylendiderin
hMPV
          APV C
APV B
          APV A
hRSV A
hrsv B
bRSV
PVM
          DEAL POLICY CONCHESTION OF LUDINGS ITKAAACYSLH
hMPV
          APV C
APV A
hrsv A
hrsv b
brsv
PVM
          NIIKQLQEVEVRQARDNKLSDSKHVALHNLVLSYMEMS-KTPASLINNLK
HMPV
          APV C
APV B
hrsv A
hRSV B
D/M
          HLPHEKLKKLAKLIIDLSAGAE--NDSSYALQDSESTNQVQ---
K.K. E. V-- TA.M. ANSD----
K. K. J.Q. S.GE.AN.NT.KGD.S.---
LIQ. P.SD.A.GNT.KGD.N.---
ADV. TI.NTL.IHKSITIN.PKESTVS.TNDHAKNNDTT-
ADV. TI.NTL.IHKSITIS.PKESTVJ.ONDQTKNNDITG
ADV. TI.NTL.IHKSITIS.PKESTVJ.ONDQTKNNDITG
.ADV. TI.NT.IHNEINGN.QGDIIVNEQNE------
VGV.CN.IQSV.SIEEKINSSMKTE-----
           RLPREKLKKLAKLIIDLSAGAE--NDSSYALQDSESTNQVQ---
MVV
APV C
APV B
APV A
hRSV A
brsv
В
                  -----EHGPVFITIEVDDM
HMPV
            APV C
APV B
APV A
 hrsv A
            MIKMTKPKIMILEDKY..SISSI..SSESMIATFINKNILQ.NHNHL.NH
MIKMTNNSNILIFPEKY..SISSL...NENDVIVLSHQNVLDYLQFQYPCN
------MIRLPKYY.AILH.MY.IRV-----NRNLTYDGSGPSTI
hRSV B
 brev
 PVM
             IWTHKDLKEAL---SDGIVKSHTNIYNCYLENIEIIYVKAYLS-
 hmpv
            IWTHKDLKEAL---SDGIVKSHTNIYNCYLENIEIIYVKAYLS---
V. KNE.VDII---TE...V.A..FK.R..D.....TF..---
INLG.I.EV---ARM.IID.I.RKQ.NECRKDFEF.AV.T.YT--
ASWS.I.EV---ANNVLID.I.RKQ.VECRKDFEFIAI.T.YN--
MYSPNQTFNEIHWT.QELIDTIQ.FLQHLGIIED.YTIYILV.---
APV C
 brsv A
            QRLLWIFDEIHWTEKNLLDATQQELQHLWIPED.YTIYILV.----
MYSQNHMLDDIYWT.QELIEDVLK.LHLSGIS.SKYVIYVLVL----
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 PVM
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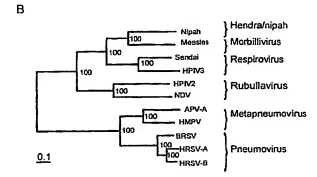


van den Hoogen et al., Figure '13



hMPV	DACTURAL SERVICE STOVENTA VOLUMENTA
APVA	RASIVTDLSKFNQAFRYETTAICADVADELHGTQSLFCWLHLIVPMTTMI
hrsvi	NCI.ISC.S.LVFAT.HV T
hrbve	NGISCSLVSTT.LV T
brsv hpiv2	
NDV	ATF.T. Q.YCLNW. Q. IKLF HATNO M LPHF E T. REMD
sv	ATF.T. Q.YCLNW.Q.IKLF.HAINQ.M.LPHF.E.IRLMDF SCFLTK.YCLNW.F.SLFGQRCN.IF.FKTF.N.M.PVLEKC.IY
hPIV3	SQFLTK.YCLNWSLFGETCNQIF.LNKNPRLEGS.TV
MV ninah	S.F.TK.YCLNWISLF.QRLN.IY.LP.F.QKRLETSVLY S.FLTKCLNWSMF.ERLIY.LPGF.N.M.KRLERSVIY
птри	R 150
<b>HMPV</b>	CAYRHAPPETKG-EYDIOKIEEQSGLYRYHMGGIEGWCOKLWTMEATSIJ
APVA hrsva	TD.GIQ.PFMTYIRDHIV.LNNVD
hrsve	
brsv	TYIRNHIT.LN.VD
PDIA5	VGDPFNAATD-AF.L., VLNGDIFIVSK
NDV	VGDPFNSDPT-DC.LSRVPNDDIYIVSARLIS.AAI
	VGDPYC.VADRM-HRQLQDHADSGIFIHNPRYLIS.AI VGDPYC.SD.E-HISLEDHPDSGFYVHNPRFLIS.AI
MV.	VSDP.CDLDA~HIPLY.VPNDCIFIK.P. v tem nv l
nipah	V.DPNCNIDK-HMELE.TP.DQIFIH.PKYSTIAT.PF.
hMPV	DVVSVKTRCQMTSLLNGDNQS1 DVSKPVKLSEG-LDEVKADYSLAVKMLK
APVA	LI.L.GKFSI.A.II.R.MQTHAQ.L.LNS. LI.L.GKFSI.A.II.R.IQTHAQ.L.LNS. LI.L.GKFSI.A.II.R.IQTHAQ.L.LNS. LI.L.GKFSI.A.II.R.IQTHAQ.L.LNS. LI.L.GKFSI.A.II.I.NQTHAQ.L.LNS. LI.L.GKFSI.A.II.I.NQTHAQ.L.LNS. LI.L.GKFSI.A.II.I.NQTHAQ.L.LNS. LI.L.GKFSI.A.I.NQTHAQ.L.LNS. LI.L.GKFSI.A.I.NQTHAQ.L.LNS. LI.L.GKFSI.A.I.NQTHAQ.L.LNS.
hrsva	.LI.L.GKFSI.A.IIR.MQTHAQLLNS
breve	LI.L.GKFSI.A.IIR.IQTHAQL.LNS
hPIV2	ILS.AESKTRVM.MVOA A TTR PR IDSTOVETTA DAGVIDER
NDV	QLAAARSH.RVACMVQV.A.TRE.RSDDSPEMVLTQLHQASDNFF.
SV	HLAA.RVGVRVSAMVQA.A.TSR.PVAQTYKQKKNHV.EEITRYFG
MA UNIAR	YLAAYESGUBIA VO TA P. DETURNING YOUR REFED
nipah	QLAAARSH.RVACMVQV A.TER.BSDDSPEMVLTQLHQASDNFF. HLAA.RVGVRVSAMVQA.TER.FVAQTYKQKKNHV.EETTRYFG HLAA.RIGVRV.AMVQA.A.TTR.FNNYDYRIKKEIV.KDV.RFFD YLAAYESGVRIAVQT.A.T.R.PSTWPYNLKKREAARVTRDYFV FLSAYE.NTRIAMIVQE.AITQK.HPNLPYKVKKEICAKQ.QLYFE
	D 250
APVA	EIRDAYRNIGHKLKEGETYISRDLQFISKVIQSEGVMHPTPIKKIRVGP AV. Y. W. T. Y. AAV. LLYKE.AG. GT. M. M. T. HN. YY. ASV. LLYKE.AG. GT. M. M. T. HN. YY. ASV. LLYKE.AS. LY M. M. T. HN. YY. ASV. LLYKE.AS. LY M. M. T. HN. YY. ASV. LLYKE.AS. GT. M. M. T. HN. YY. ASV. LLYKE.AG. GT. M. M
hrsva	LLYKE.AGGT M. M. T. HN. YY.AS. V
hrsvb	LLYKE.AGGTMMTHNYY.ASV
hPIV2	BL ANNYGL O AO I STEELY DISTO DISTON
NDV	LIHVNHLNDRIR.DTFFIYR.FKD.AILSOVL.NSSKIVI
sv	AL. HVMFD E LN I SKMFVY . R . YYD . KIL . QCL . ALT . CVF
PLIA3	SL.EVMDDL.E.LN.I.SKMFIY.R.YYD.RIL.QAL.ALS.CVF
nipah	RL.MNL.ALNATITH.FIYK.HYD.AVISOAL SWI CCE
	300
hmpv Apva	WINTILDDIKTSAESIGSLCQELEFRGESIIVSLILRNFWLYNLYMHESK
brsva	F.V.L. T. V. IIC F W OTHER
THICO A D	A CONTRACTOR OF TOTAL OF THE CONTRACTOR OF THE C
PDIM	TROUT CECTOR CONCERNMENTS THE
MDA	VSGDLSENTVM.CAN.A.TVAR CEN BEAGG V INVEN G
24	.SE.LV.ENRSACSN.STSIAKAIENYSDIIC CIALVETC
MA.	SE.VI.ETRSASSNLATSFAKAIENYSPVIC ACSIEKNI
	.SEV.ETRAACSN.ATTMAKSIEYDRYLA.S.NVLKVI .SE.LV.ETRSACSN.STTIAKAIENSRNVG.CINILKVI
	279
hMPV	QHPLAGKQLFKQLNKTLTSVQRFFEIKKE
APVA hrsva	D.SA.ISNERK.KRLAD. N.A.CNNK.YLDIL.V.KHLKTNLDNI
hrsvb	N.A.CNNK.YLDIL.V.KHLKT., NLDST
brsv	N.A.CHNK.YLDIL.V.AHLKM., NLDNT
hPIV2	RQLTYDLIFFQYSIPGE.ISEI.LQHPRL
NDV SV	VQTYFDSEFSITN.SHPDLN.SWI.DISF .QVCISLGMTINPTISP.VRDQY.KG.NW
	.QLYIALGMNINPTI.QNIKDQY.KNSNW
MA	.QI.ISLGFTINSTM.RDV.IPLLTNNDL
nipah	.QL.ISTEFSINETLDVTSPISNNLDW





Α						
	END	SPACER	START			
N			GGGACAAGUGAAAAUG			
N/P	GAGUAA	(N)10	GGGACAAGUGAAAAUG			
P/M	AUGUAG	(N) 19	GGGACAAGUCAAAAUG			
M/F	AGAUAA	(N) 109	GGGACAAAUAAAAAUG			
F/M2	AAUUAG	(N) 52	GGGACAAAUCAUAAUG			
M2/SH	AGUUAG	(N) 17	GGGAUAAAUGACAAUG			
SH/G	UCCUAA	(N) 188	GGGACAAGUAGUUAUG			
G/L	AGU <u>UAA</u>	(N)196	GAGACAAAUAACAAUG			
L	AAA <u>UGA</u>					
В						
hMPV Le: 3'UGCUCUUUUUUUGCGCAUAUUUAAUCUAAGGUUUUUUUUAUA						
hMPV Tr: 5'ACGAGAAAAAAC CGUAUACAUCCAAUUAUAAUUUCUUAU						
hmpv Tr: 5'ACGAGAAAAAACCGUAUACAUCCAAUUAUAAUUUCUUAUUUUUA						
	11111		[ [ [ ] ] ] ] ] [ ] [ ] [ ] [ ] [ ] [ ]			
APV T	r: 5'ACGAGA	AAAAAACGCUAUUCAU	AAAUUUUUAGCUUUUAGUUUUU			
hMPV L	e: 3'UGCUCU	UUUUUUGCGCAYAUUU	AUCUAAGGUUUUUUUAU AEGG			
	111111		11 111 1 11 111 1111			
APV L	illill e: 3'UGCUCU					
Arv L	e: 3.00COCO	UUUUUUGCGUA~AGUU(	CGUCCAAGAUCUUUUUUUUUAUUAEGQ			